US ERA ARCHIVE DOCUMENT



# Reregistration Eligibility Decision (RED)

Cedarwood Oil







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### GLOSSARY OF TERMS AND ABBREVIATIONS

a.i. Active Ingredient

CAS Chemical Abstracts Service

CSF Confidential Statement of Formula

EEC Estimated Environmental Concentration. The estimated pesticide concentration

in an environment, such as a terrestrial ecosystem.

EP End-Use Product

EPA U.S. Environmental Protection Agency

FIFRA Federal Insecticide, Fungicide, and Rodenticide Act

FFDCA Federal Food, Drug, and Cosmetic Act

HDT Highest Dose Tested

LC<sub>50</sub> Median Lethal Concentration. A statistically derived concentration of a substance

that can be expected to cause death in 50% of test animals. It is usually expressed as the weight of substance per weight or volume of water or feed, e.g.,

mg/l or ppm.

LD<sub>50</sub> Median Lethal Dose. A statistically derived single dose that can be expected to

cause death in 50% of the test animals when administered by the route indicated (oral, dermal). It is expressed as a weight of substance per unit weight of animal,

e.g., mg/kg.

LD<sub>lo</sub> Lethal Dose-low. Lowest Dose at which lethality occurs

LEL Lowest Effect Level

LOEL Lowest Observed Effect Level

MP Manufacturing-Use Product

MPI Maximum Permissible Intake

MRID Master Record Identification (number). EPA's system of recording and tracking

studies submitted.



## GLOSSARY OF TERMS AND ABBREVIATIONS (cont.)

N/A Not Applicable

NPDES National Pollutant Discharge Elimination System

NOEL No Observed Effect Level

OPP Office of Pesticide Programs

PADI Provisional Acceptable Daily Intake

ppm Parts Per Million

RfD Reference Dose

RS Registration Standard

TD Toxic Dose. The dose at which a substance produces a toxic effect.

TC Toxic Concentration. The dose at which a substance produces a toxic effect.

TMRC Theoretical Maximum Residue Contribution.

### EXECUTIVE SUMMARY

The Agency has completed its reregistration assessment of the available information on the pesticide active ingredient cedarwood oil in the case named Wood Oils and Gums. It has been determined that the currently registered uses will not cause unreasonable risk to humans or the environment and these uses are eligible for reregistration.

Cedarwood oil is a natural component of wood from the tree, <u>Juniperus virginiana</u> L., for use in pesticide products to repel moths from and inhibit mildew in clothing; and repel fleas from pets and their sleeping quarters. Current registered products include cedarwood blocks, a pet collar, and a ready-to-use liquid. The Agency is proposing, under a separate action, to deregulate the cedarwood block products in accordance with FIFRA 25(b).

Before reregistering the products containing cedarwood oil, the Agency is requiring that additional technical chemistry data on the extracted oil, as well as product specific data on acute toxicology, chemistry and efficacy, revised Confidential Statements of Formula (CSF) and revised labeling be submitted within eight months of the issuance of this document. After reviewing these data and revised labels and finding them acceptable in accordance with Section 3(c)(5) of FIFRA, the Agency will reregister products. Those products which contain other active ingredients will be eligible for reregistration only when the other active ingredients are determined to be eligible for reregistration.



### I. INTRODUCTION

In 1988, the Federal Insecticide, Fungicide, and Rodenticide Act (FIFRA) was amended to accelerate the reregistration of products with active ingredients registered prior to November 1, 1984. The amended Act provides a schedule for the reregistration process to be completed in nine years. There are five phases to the reregistration process. The first four phases of the process focus on identification of data requirements to support the reregistration of an active ingredient and the generation and submission of data to fulfill the requirements. The fifth phase is a review by the U.S. Environmental Protection Agency (referred to as "the Agency") of all data submitted to support reregistration.

FIFRA Section 4(g)(2)(A) states that in Phase 5 "the Administrator shall determine whether pesticides containing such active ingredient are eligible for reregistration" before calling in data on products and either reregistering products or taking other "appropriate regulatory action." Thus, reregistration involves a thorough review of the scientific data base underlying a pesticide's registration. The purpose of the Agency's review is to reassess the potential hazards arising from the currently registered uses of the pesticide; to determine the need for additional data on health and environmental effects; and to determine whether the pesticide meets the "no unreasonable adverse effects" criterion of FIFRA.

This document presents the Agency's decision regarding the reregistration eligibility of the registered uses of cedarwood oil. The document consists of six sections. Section I is the introduction. Section II describes cedarwood oil, its uses, data requirements and regulatory history. Section III discusses the human health and environmental assessment based on the data available to the Agency. Section IV presents the reregistration decision for cedarwood oil. Section V discusses the reregistration requirements for cedarwood oil. Finally, Section VI is the Appendices which support this Reregistration Eligibility Decision document. Additional details concerning the Agency's review of applicable data are available on request.<sup>1</sup>



<sup>&</sup>lt;sup>1</sup>EPA's reviews of data on the set of registered uses considered for EPA's analysis may be obtained from the OPP Public Docket, Field Operations Division (H7506C), Office of Pesticide Programs, EPA, Washington, DC 20460.

### II. CASE OVERVIEW

### A. Chemical Overview

The following active ingredient is covered by this Reregistration Eligibility Decision document:

• Common Name: Cedarwood oil

• Chemical Name: Oil extracted from species of cedar trees, especially

Juniperus virginiana L. (1)

• Chemical Family: Wood oils and gums

• CAS Registry Number: 800-27-9

• OPP Chemical Code: 40505

• Empirical Formula: cedarwood oil is a mixture of organic compounds

• Trade and Other Names: oil of cedar

• Basic Manufacturer: not applicable

### B. Use Profile

The following is information on the current registered uses with an overview of use sites and application methods. Appendix A is a detailed table of these uses of cedarwood oil.

### For Cedarwood oil:

Type of Pesticide: repellant/feeding depressant; fungicide

Use Sites: Indoor Residential - domestic dwellings and their contents,

Pets/Animals (presumably cats and dogs) and their living and sleeping
quarters

Target Pests: fleas, moths, mildew

### Formulation Types Registered:

Liquid ready-to-use - sprayed on animal bedding

0.48 % cedarwood oil

### Impregnated pet collar/tag

0.5 % oil of citronella 1.0 % oil of eucalyptus 0.5 % cedarwood oil 2.0 % oil of pennyroyal 0.125 % oils, rue;

Wood blocks, containing an average of --

5.17% cedarwood oil (2-8% range)

### C. Data Requirements

The Agency has waived all generic data requirements except for physical chemistry for this active ingredient. The rationale for this action is discussed below and also in Section III. Instead it has relied on general, commonly available information about cedarwood oil. Appendix B includes all data requirements identified by the Agency for currently registered uses needed to support reregistration.

EPA has developed a target database, set forth in the regulations (2) and the Agency's Reregistration Phase 2 Technical Guidance Document, to be addressed for pesticide reregistration. These regulations and the Guidance Document specify the necessary data based on factors including use sites, potential environmental and human (dietary and occupational) exposures, product formulation types, and product application methods. Due to the-diverse-nature and characteristics of pesticide products and their uses subject to reregistration, the Agency also recognizes the necessity to modify the data requirements for specific pesticides, including waiving certain data requirements because such requirements are inappropriate or unnecessary for risk assessment and reregistration.



This approach to waiving individual data requirements has served to identify the appropriate data requirement sets for pesticide products. Further, the Agency believes there is a category of pesticide active ingredients for which a broadly reduced set of data requirements are appropriate for reregistration. Specifically, products in this category may be exempt from the generic data requirements for toxicology, residue chemistry, human exposure, ecological effects, and environmental fate on the active ingredient. The Agency believes there are considerations which, when taken together, can form the basis for a conclusion that such a reduction in data requirements is appropriate for a particular pesticide active ingredient, while not compromising human health or environmental safety.

There are, however, certain data requirements which are essential and not likely to be waived. Basic product identity/chemistry information on the active ingredient and formulated products is required for pesticides in this category so that the Agency has reasonable certainty of the pesticide's chemical and physical characteristics. Also, product specific acute toxicology studies are required for the Agency to determine appropriate product labeling for potential hazards to those who handle or apply these products. However, these toxicology studies may also be waived if an assessment of the product formulation, including the inert ingredients, indicates that such studies are unnecessary to prescribe appropriate labeling. Efficacy studies may be required of formulated products labeled for uses and pests that are of a public health concern.

In considering cedarwood oil for reregistration eligibility the Agency believes it is an active ingredient that should be considered for this broad waiver of the generic data requirements. The considerations that led the Agency to this conclusion are discussed in Section III below.

### D. Regulatory History

Originally, the reregistration case Wood Oil and Gums included three active ingredients: cedarwood oil, canadian balsam, and ester gum. After Phase 2 in 1990, products containing the latter two active ingredients were cancelled for non-support by the registrants, leaving only products containing cedarwood oil.

Cedarwood oil was initially registered as a pesticide in the United States in 1960 to repel moths from clothing. Since then products have been registered to repel fleas from pets and their bedding. The five currently registered products containing cedarwood oil as an-active-ingredient fall under one of three types of pesticide products: three are solid cedarwood block products, ("Cedar Fresh", EPA Reg. No. 65555-1; "Ozark Cedar", EPA Reg. No. 65813-1, and "Woodland/RPM Cedar", EPA Reg. No. 66211-1), one product is a pet collar ("Herbal Flea Collar", EPA Reg. No. 42443-1), and one product is a ready-to-use liquid that may be applied by hand spray ("Green Earth

### III. SCIENCE ASSESSMENT

### A. Physical Chemistry Assessment

Cedarwood oil is a distilled extract from the cured cedarwood obtained from <u>Juniperus virginiana</u> L., and other species of cedar. The chief chemical components are cedrene (a terpene) and cedral (cedar camphor).

Cedarwood oil ranges in color from colorless to slightly yellow. It is somewhat viscid. It is insoluble in water but soluble in both alcohol and ether (1).

### B. Human Health And Environmental Assessment

As discussed above, the Agency has waived the generic data requirements, except for certain technical chemistry information, for cedarwood oil. It is relying on commonly available information about this chemical and its uses to reach a decision about its potential risks to human health and the environment associated with the current uses of registered products. The specific information about cedarwood oil the Agency considered is as follows:

Cedarwood oil is a mixture of organic compounds. It is a component of many non-pesticidal consumer products currently marketed in the United States. Cedarwood oil is listed as a food additive by the Food and Drug Administration (5). The alcohols and terpenes of cedarwood oil are considered by FDA to be Generally Recognized As Safe (GRAS). As a pesticide, it repels insect pests by a non-toxic mode of action. Its mode of action against mildew is unknown. EPA is not aware of any adverse effects of the active ingredient to humans or the environment in the literature when used in a manner prescribed in end-use product labeling. There have been no reported incidents of toxicity.

There have been some studies performed on cedarwood dust as well as the constituents of cedarwood oil. According to a recent survey of the literature, pulmonary effects and liver effects have been noted in laboratory animals (3). These effects are hypothesized to be related to the occupational hazards associated with saw mill workers (4) chronically exposed to environments high in cedarwood dust. However, this type of exposure is not indicative of that from use of the currently registered pesticide products.

The Agency believes there is negligible human and environmental exposure to the pesticide as a result of the use patterns; there is a low use rate and frequency of



application and/or the products are applied in a confined or contained manner. However, the handling and use of the liquid product could pose a relatively greater exposure potential by the dermal and inhalation routes. Product specific acute toxicity testing will allow the Agency to address appropriate labeling to address potential concerns for users.

Since the pesticide will be used in indoor domestic dwellings, on pets and on their living and sleeping quarters, the Agency expects that there will be negligible exposure to the environment and to nontarget organisms.

Based on these factors the Agency does not believe generic data, beyond those data required to satisfy basic characterization of the chemistry (refer to Appendix B), are necessary to determine whether the current registered uses of this active ingredient pose unreasonable risks to humans or the environment. However, it is requiring the submission of product specific data (chemistry, acute toxicity and efficacy).

In conclusion, the Agency has determined that the use of cedarwood oil as an active ingredient in products for the current uses should not result in unreasonable adverse effects to human health or the environment.

### IV. RISK MANAGEMENT AND REREGISTRATION ELIGIBILITY DECISION

### A. Determination of Reregistration Eligibility

Section 4(g)(2)(A) of FIFRA calls for the Agency to determine, after submission of relevant data concerning an active ingredient, whether products containing the active ingredients are eligible for reregistration. As discussed above, the Agency has determined that the set of generic data requirements that would normally be applicable to cedarwood oil need not be satisfied for the Agency to reach a decision on potential risks and reregistration eligibility. Rather, it has considered general and commonly available information. The Agency has determined that cedarwood oil meets criteria as outlined in the document "Guidance for Making Determinations to Reduce Data Requirements". Cedarwood oil met the criteria due to its use and availability for nonpesticide food uses, its regulatory status as a chemical classified as GRAS and exempt from the requirement of food additive tolerances; its non-toxic mode of action as a pesticide; that there is negligible human and environmental exposure to the pesticide as a result of the porposed use pattern, and the lack of reports of adverse effects. (No data were submitted-under-6(a)(2) of FIFRA, no significant incidents have been reported to the Agency, and there is no indication in the literature that the pesticide poses adverse effects in humans or to the environment when used in a manner prescribed in end-use poduct labeling.) Appendix B identifies the sources for this information that the Agency reviewed as part of its determination of reregistration eligibility of cedarwood oil and

lists the submitted studies that the Agency considered acceptable.

The Agency believes this information is sufficient to support reregistration and, that cedarwood oil can be used without resulting in unreasonable adverse effects to human health and the environment. The Agency therefore finds that all products containing cedarwood oil as the active ingredient are eligible for reregistration. The reregistration of particular products is addressed in Section V of this document.

Although the Agency has found that all uses of cedarwood oil are eligible for reregistration, it should be understood that the Agency may take appropriate regulatory action, and/or require the submission of additional data to support the registration of products containing cedarwood oil, if new information comes to the Agency's attention or if the data requirements for registration (or the guidelines for generating such data) change.

However, the Agency has determined that natural cedarwood products consisting of wood blocks, not treated or impregnated with any additional substance(s), distributed and sold as moth and flea repellents or mildew control agents, are of a character which is unnecessary to be subject to FIFRA. Cedarwood oil is naturally contained in the cedar wood; it cannot easily be separated from the wood or wood products. Consumers using these cedarwood products are unlikely to be exposed to significant amounts of cedarwood dust or oil, either by the inhalation or dermal route. The Agency believes there is negligible risk to man or the environment associated with the use of these wood products as pesticides. Therefore, under authority of FIFRA section 25(b), 7 U.S.C 136w(b), the Agency has proposed that such products be exempted from provisions of FIFRA, except misbranding provisions (58 FR 42711; August 11, 1993). A 30-day comment period is provided. Products containing cedarwood oil extracted from wood or synthesized and subsequently used for pesticide purposes are not included in this proposed exemption.

Since the Agency is proposing that these cedarwood products be exempted from regulation, further reregistration requirements are being held in abeyance. If the Agency concludes it should not proceed to effect the exemption of these products, the Agency will proceed to impose the appropriate reregistration requirements on these products.

### 1. Eligible and Ineligible Uses

The Agency has determined that all current uses of cedarwood oil products are eligible for reregistration.

### B. Risk Management Decision

In consideration of the above information about cedarwood oil, the Agency finds no reason to impose new risk reduction measures for currently registered uses. The



Agency will however, assess the need for product specific risk reduction measures upon receipt of data that are being required under the Product Specific Data Call-in Notice appended to this document.

### V. ACTIONS REQUIRED BY REGISTRANTS

This section specifies the data requirements and responses necessary for the reregistration of end-use products. Because there are no registered manufacturing-use products or technical products the generic data requirements are required for all registrants.

### A. Technical Grade Information

### 1. Generic Data Requirements

There are currently no registered manufacturing use products for cedarwood oil. The generic data base supporting the reregistration of products containing cedarwood oil for the above eligible uses has been reviewed and determined to be incomplete. Registrants are required to submit the technical chemistry data corresponding to Series 61 and Series 62 for the analysis and certification of product ingredients. If the product is a United States Pharmacopoeia (USP) grade, a copy of USP analysis with citation of the analytical methods used and certification would satisfy the requirement for Series 62.

The Confidential Statement of Formula (CSF) must be supported by analytical data. The data on the physical and chemical characteristics of cedarwood oil from the Material Safety Data Sheet (MSDS) for the product may be compiled by the registrant in the format required by the FIFRA Accelerated Reregistration Phase 3 Technical Guidedance, specifically PR Notice 86-5 to satisfy some of the requirements of Series 63. The generic data requirements are listed in Appendix F, the Generic Data Call-in Notice.

### **B.** End-Use Products

### 1. Additional Product-Specific Data Requirements

Section 4(g)(2)B) of FIFRA calls for the Agency to obtain any needed product-specific data regarding the pesticide after a determination of eligibility has been made. The product specific data requirements are listed in Appendix G, the Product Specific Data Call-In Notice.

Registrants must review previous data submissions to ensure that they meet current EPA acceptance criteria (Appendix F; Attachment E) and if not, commit to conduct new studies. If a registrant believes that previously submitted data meet current testing standards, then study MRID numbers should be cited according to the instructions in the Requirement Status and Registrants Response Form provided for each product.

Additional physical chemistry data on cedarwood oil are required to provide the Agency with a more complete characterization of the chemistry of the cedarwood oil that is used in the registered products. These generic data requirements are imposed on the end-use products because there is no registered source of technical grade cedarwood oil.

### 2. Labeling Requirements for End-Use Products

The labels and labeling of all products must comply with EPA's current regulations and requirements as specified in 40 CFR §156.10. Please follow the instructions in the Pesticide Reregistration Handbook with respect to labels and labeling. In addition, registrants with products that are used on pets and/or animals will be required to specify on their labels which animals may be treated with the product.

### A. Existing Stocks

Registrants may generally distribute and sell products bearing old labels/labeling for 26 months from the date of the issuance of this Reregistration Eligibility Decision document (RED). Persons other than the registrant may generally distribute or sell such products for 50 months from the date of the issuance of the RED. However, existing stocks time frames will be established case-by-case, depending on the number of products involved, the number of label changes, and other factors. Refer to "Existing Stocks of Pesticide Products; Statement of Policy"; Federal Register, Volume 56, No. 123, June 26, 1991.

The Agency has determined that registrants may distribute and sell cedarwood oil products bearing old labels/labeling for 26 months from the date of issuance of this RED. Persons other than the registrant may distribute or sell such products for 50 months from the date of the issuance of this RED.



# VI. APPENDICES

# APPENDIX A

**Table of Use Patterns Subject to Reregistration** 

Page 1	
	Use Limitation Codes
NO.	Geographic Dissilowed
mical 040505 [Cedarwood oil]	ifn. Restr. Geographic Interv Entry Allowed (days) Interv (days)
[Wood oils and game] Chemical 040505	Mex. Min Appe Int 8 Mex (de Rate
- CASE 3150,	Maximum Soil Max Application Text # Rates (Max Apps Dse)
APPENDIX A	Form Minimum Application Rate
Date 09/15/93 - Time 15:39	SITE Application Type, Application Timing, Application Equipment Surface Type & Efficacy Influen- cing Factor (Antimicrobial only)

USES ELIGIBLE FOR REREGISTRATION

NON-FOOD/NON-FEED

		2		*		<b>*</b>	¥		2	*
		2		S		₹	2		₹	2
	USE Group: INDOOR RESIDENTIAL	S	Use Group: INDOOR RESIDENTIAL	<u>\$</u>	Use Group: INDOOR RESIDENTIAL	<b>£</b>	2	Use Group: INDOOR RESIDENTIAL	<b>S</b>	**
		HA . HS	INDOOR	NA * NS	THOOOR	* * *	₩.	1MDOOM1	MA * HS	<u>*</u>
	ŝ	•	ğ		ë	•	*	ÿ	•	•
•	13 es0	¥	Use Gr	¥	Use Gr	¥	¥	Use Gr	¥	.0003125 (b •
		IMPR NA		INPR NA		¥	<b>4</b>		¥	<b>\$</b>
		Ĭ		<u> </u>		\$	<u>=</u>		₹	15.
PARKETERS PARKETT PLET THE STATES	SECTION POST OF PRECIOUS CONTENTS	Funigation., When needed., By hard.	HOUSEHOLD/DOMESTIC DWELLINGS INDOOR PRENISES	Funigation., When needed., By hand.	PET LIVING/SLEEPING QUARTERS	Animal bedding treatment., When needed., Pump spray bottle.	Fumigation, When needed, By hand.	SITE TERM TOO GENERAL	Animal treatment (spray)., When needed., Pump spray bottle.	Flee coller, When needed, By hand.

LEGEND

```
HEADER ABBREVIATIONS
HAX. # Apps
HAX. # Apps
HAX. Apps
HAX. Apps B MAX Rate : Maximum number of Applications at Maximum Dosage Rate
Min. Interv (days) : Minimum Interval between Applications (days)
Restr. Entry Interv (days): Restricted Entry Interval (days)
```

SOIL TEXTURE FOR MAX APP. RATE

Coarse

Fine

: Others

FORMULATION CODES
1C/T : IMPREGNATED COLLAR/TAG
1MPR : IMPREGNATED MATERIAL
RTU : LIQUID-READY TO USE

ABREVIATIONS
AN : As Needed
NA : Not Applicable
NS : Not Specified (on Label)

APPLICATION RATE
DCNC : Dosage Can Not be Calculated
No Calc : No Calculation can be made
W : PPM Calculated by weight
Y : PPM Calculated by yolume
Cut : Mundred Weight

16

Table of the Generic Data Requirements and Studies Used to Make the Reregistration Decision

### GUIDE TO APPENDIX B

Appendix B contains listings of data requirements which support the reregistration for the pesticide bromine covered by this Reregistration Eligibility Decision. It contains generic data requirements that apply to bromine in all products, including data requirements for which a "typical formulation" is the test substance.

The data table is organized in the following format:

- 1. <u>Data Requirement</u> (Column 1). The data requirements are listed in the order in which they appear in 40 CFR, Part 158. The reference numbers accompanying each test refer to the test protocols set in the Pesticide Assessment Guidelines, which are available from the National Technical Information Service, 5285 Port Royal Road, Springfield, VA 22161 (703) 487 4650.
- 2. <u>Use Pattern</u> (Column 2). This column indicates the use patterns for which the data requirements apply. The following letter designations are used for the given use patterns:
  - A Terrestrial food
  - B Terrestrial feed
  - C Terrestrial non-food
  - D Aquatic food
  - E Aquatic non-food outdoor
  - F Aquatic non-food industrial
  - G Aquatic non-food residential
  - H Greenhouse food
  - I Greenhouse non-food
  - J Forestry
  - K Residential
  - L Indoor food
  - M Indoor non-food
  - N Indoor medical
  - O Indoor residential
- 3. <u>Bibliographic citation</u> (Column 3). If the Agency has acceptable data in its files, this column lists the identifying number of each study. This normally is the Master Record Identification (MRID) number, but may be a "GS" number if no MRID number has been assigned. Refer to the Bibliography appendix for a complete citation of the study.



Data Supporting Guideline Requirements for the Reregistration of Cedarwood Oil

REQUI	REQUIREMENT	USE PATTERN	CITATION	
PROD	PRODUCT CHEMISTRY		-	
61-1	Chemical Identity	All	42319001, 42101201, 41549503	
61-2A	Start. Mat. & Mnfg. Process	ΑШ	42101201, 42362601, 42319001	
61-2B	Formation of Impurities	All	42101201, 42362601, 42319001	
62-1	Preliminary Analysis	All	42362602, 42319002	
62-2	Certification of limits	All	42362602, 42319002	
62-3	Analytical Method	All	42319002	
63-2	Color	All	SATISFIED	
63-3	Physical State	All	SATISFIED	
63-4	Odor	All	SATISFIED	
63-5	Melting Point	All	SATISFIED	
63-6	Boiling Point	All	SATISFIED	
63-7	Density	All	SATISFIED	
63-8	Solubility	All	SATISFIED	,
63-9	Vapor Pressure	All	SATISFIED	. *
63-10	Dissociation Constant	ALL	SATISFIED	
63-11	Octanol/Water Partition	ALL	SATISFIED	
63-12	Hď	ALL	SATISFIED	÷

Data Supporting Guideline Requirements for the Reregistration of Cedarwood Oil

REQUIREMEN	REQUIREMENT	USE PATTERN	CITATION
PROD	PRODUCT CHEMISTRY		
63-131	Stability	ALL	SATISFIED
ECOL	ECOLOGICAL EFFECTS		
71-1A	Acute Avian Oral - Quail/Duck	ALL	WAIVED
71-2A	Avian Dietary - Quail	ALL	WAIVED
72-1A	Fish Toxicity Bluegill	ALL	WAIVED
72-1C	Fish Toxicity Rainbow Trout	ALL	WAIVED
72-2A	Invertebrate Toxicity	ALL	WAIVED
TOXIC	TOXICOLOGY		
81-1	Acute Oral Toxicity - Rat	ALL	WAIVED
81-2	Acute Dermal Toxicity - Rabbit/Rat	ALL	WAIVED
81-3	Acute Inhalation Toxicity - Rat	ALL	WAIVED
81-4	Primary Eye Irritation - Rabbit	ALL	WAIVED
81-5	Primary Dermal Irritation - Rabbit	ALL	WAIVED
81-6	Dermal Sensitization - Guinea Pig	ALL	WAIVED



<sup>1</sup> Confirmatory data are being required from certain registrants

-	Data Supporting Guideli	ne Kequirem	Data Supporting Guideline Requirements for the Reregistration of Cedarwood Oil
REQUI	REQUIREMENT ,	USE PATTERN CITATION	CITATION
TOXI	TOXICOLOGY		
84-2A	84-2A Gene Muatation-(Ames Test)	All	WAIVED
84-2B	Structural Chromosomal Aberration	ALL	WAIVED
84-4	Other Genotoxic Effects	ALL	WAIVED

	Data	Data Supporting	Guideline Requi	reme	Guideline Requirements for the Reregistration of Cedarwood On
REQUIREMENT	EQUIREMENT	-	USE PATTI	RN	USE PATTERN CITATION
ENVIR	ONMEN	ENVIRONMENTAL FATE		•	
161-1 Hydrolysis	Hydrolysi	. <b>s</b> a	ALL		WAIVED



## **APPENDIX C**

Citations Considered to be Part of the Data Base Supporting the Reregistration of Wood Oils and Gums

#### GUIDE TO APPENDIX C

- 1. CONTENTS OF BIBLIOGRAPHY. This bibliography contains citations of all studies considered relevant by EPA in arriving at the positions and conclusions stated elsewhere in the Reregistration Eligibility Decision. Primary sources for studies in this bibliography have been the body of data submitted to EPA and its predecessor agencies in support of past regulatory decisions. Selections from other sources including the published literature, in those instances where they have been considered, are included.
- 2. UNITS OF ENTRY. The unit of entry in this bibliography is called a "study". In the case of published materials, this corresponds closely to an article. In the case of unpublished materials submitted to the Agency, the Agency has sought to identify documents at a level parallel to the published article from within the typically larger volumes in which they were submitted. The resulting "studies" generally have a distinct title (or at least a single subject), can stand alone for purposes of review and can be described with a conventional bibliographic citation. The Agency has also attempted to unite basic documents and commentaries upon them, treating them as a single study.
- 3. IDENTIFICATION OF ENTRIES. The entries in this bibliography are sorted numerically by Master Record Identifier, or "MRID number". This number is unique to the citation, and should be used whenever a specific reference is required. It is not related to the six-digit "Accession Number" which has been used to identify volumes of submitted studies (see paragraph 4(d)(4) below for further explanation). In a few cases, entries added to the bibliography late in the review may be preceded by a nine character temporary identifier. These entries are listed after all MRID entries. This temporary identifying number is also to be used whenever specific reference is needed.
- 4. FORM OF ENTRY. In addition to the Master Record Identifier (MRID), each entry consists of a citation containing standard elements followed, in the case of material submitted to EPA, by a description of the earliest known submission. Bibliographic conventions used reflect the standard of the American National Standards Institute (ANSI), expanded to provide for certain special needs.
  - Author. Whenever the author could confidently be identified, the Agency has chosen to show a personal author. When no individual was identified, the Agency has shown an identifiable laboratory or testing facility as the author. When no author or laboratory could be identified, the Agency has shown the first submitter as the author.
  - b. Document date. The date of the study is taken directly from the document. When the date is followed by a question mark, the bibliographer has deduced the date from the evidence contained in the document. When the date appears as (19??), the Agency was unable to determine or estimate the date of the document.



- c. Title. In some cases, it has been necessary for the Agency bibliographers to create or enhance a document title. Any such editorial insertions are contained between square brackets.
- d. Trailing parentheses. For studies submitted to the Agency in the past, the trailing parentheses include (in addition to any self-explanatory text) the following elements describing the earliest known submission:
  - (1) Submission date. The date of the earliest known submission appears immediately following the word "received."
  - (2) Administrative number. The next element immediately following the word "under" is the registration number, experimental use permit number, petition number, or other administrative number associated with the earliest known submission.
  - (3) Submitter. The third element is the submitter. When authorship is defaulted to the submitter, this element is omitted.
  - (4) Volume Identification (Accession Numbers). The final element in the trailing parentheses identifies the EPA accession number of the volume in which the original submission of the study appears. The six-digit accession number follows the symbol "CDL," which stands for "Company Data Library." This accession number is in turn followed by an alphabetic suffix which shows the relative position of the study within the volume.

- The Merck Index; An Encyclopedia of Chemicals, Drugs, and Biologicals. Windholz, Martha, editor, et. al. Tenth Edition. Published in 1983 by Merck and Company, Rahway New Jersey, U.S.A.
   Code of Federal Regulations, Title 40, part 158, revised as of July 1, 1992. Published by the Office of the Federal Register National Archives and Records Administration, Washington, D.C., U.S.A.
   Gordon WP; Forte AJ; McMurtry RJ; Nelson SD (1982) Hepatotoxicity and Pulmonary Toxicity of Pennyroyal Oil and its Constituent Terpenes in the Mouse. Published by the Journal of Toxicology and Applied Pharmacology; Volume 65, ISS 4, P413-24.}
   Ayars GH; Altman LC; Frazier CE; Chi EY; (1989) The Toxicity of
  - Constituents of Cedar and Pine Woods to Pulmonary Epithelium. Published by Journal of Allergy and Clinical Immunology; Volume 83, ISS 3, P610.
- 5. Code of Federal Regulations, Title 21, part 172, section 515, revised as of July 1, 1992. Published by the Office of the Federal Register National Archives and Records Administration, Washington, D.C., U.S.A.
- 41549503 Gunther (19??) Oil of Cedarwood. pp. 356-363 in unknown source.
- 42101201 Arbor American Corp. (1991) Chemical Identity: Cedar Wood Oil. Unpublished study. 6 p.
- Anon. (1992) Product Identification and Disclosure of Ingredients; Description of Beginning Materials and Manufacturing process; Discussion of formation of impurities (contains published material). Unpublished study prepared by Cor-Pak International. 15 p.
- 42319002 Maltese, L. (1992) Analysis of Redwood Cedars: Lab Project Number: 92-1076: 92-1117: 92-1216. Unpublished study prepared by Stillwell & Gladding. 13 p.
- Anon. (1992) Description of Materials Used to Produce Product and Description of Production Process: Aromatic Cedar. Unpublished study prepared by P & M —Consumer Products, Inc. 7 p.
- 42362602 Zavarin, E. (1992) Analysis of Red Cedar Wood Juniperus Virginiana: Unpublished study prepared by Forest Products Laboratory (University of California at Berkley). 8 p.



# APPENDIX D List of Available Related Documents

The following is a list of available documents related to cedarwood oil. It's purpose is to provide a path to more detailed information if it is needed. These accompanying documents are part of the Administrative Record for cedarwood oil and are included in the EPA's Office of Pesticide Programs Public Docket.

- 1. Health and Environmental Effects Science Chapters
- 2. Detailed Label Usage Information System (LUIS) Report
- 3. Cedarwood oil RED Fact Sheet
- 4. PR Notice 91-2 (included in this appendix) pertains to the Label Ingredient Statement
- 5. Guidance for Making Determinations to Reduce Data Requirements
- 6. Minutes from the April 19 and May 17 meetings of the Ad Hoc Screening Committee for Reduced Data Requirements



## APPENDIX E

Pesticide Reregistration Handbook PR Notices 86-5 and 91-2 PESTICIDE REGISTRATION HANDBOOK

PR Notice 86-5



## UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

WASHINGTON, D.C. 20460

July 29, 1986

OFFICE OF

PR NOTICE 86-5

PREVENTION, PESTICIDES AND TOXIC SUBSTANCES

NOTICE TO PRODUCERS, FORMULATORS, DISTRIBUTORS AND REGISTRANTS

Attention:

Persons responsible for Federal registration of

pesticides.

Subject:

Standard format for data submitted under the Federal Insecticide, Fungicide, and Rodenticide Act (FIFRA) and certain provisions of the Federal

Food, Drug, and Cosmetic Act (FFDCA).

#### I. Purpose

To require data to be submitted to the Environmental Protection Agency (EPA) in a standard format. This Notice also provides additional guidance about, and illustrations of, the required formats.

#### II. Applicability

This PR Notice applies to all data that are submitted to EPA to satisfy data requirements for granting or maintaining pesticide registrations, experimental use permits, tolerances, and related approvals under certain provisions of FIFRA and FFDCA. These data are defined in FIFRA §10(d)(1). This Notice does not apply to commercial, financial, or production information, which are, and must continue to be, submitted differently under separate cover.

#### III. Effective Date

This notice is effective on November 1, 1986. Data formatted according to this notice may be submitted prior to the effective date. As of the effective date, submitted data packages that do not conform to these requirements may be returned to the submitter for necessary revision.

#### IV. Background

On September 26, 1984, EPA published proposed regulations in the Federal Register (49 FR 37956) which include Requirements for Data Submission (40 CFR §158.32), and Procedures for Claims of Confidentiality of Data (40 CFR §158.33). These regulations



specify the format for data submitted to EPA under Section 3 of FIFRA and Sections 408 and 409 of FFDCA, and procedures which must be followed to make and substantiate claims of confidentiality. No entitlements to data confidentiality are changed, either by the proposed regulation or by this notice.

OPP is making these requirements mandatory through this Notice to gain resource-saving benefits from their use before the entire proposed regulation becomes final. Adequate lead time is being provided for submitters to comply with the new requirements.

## V. Relationship of this Notice to Other OPP Policy and Guidance

While this Notice contains requirements for organizing and formatting submittals of supporting data, it does not address the substance of test reports themselves. "Data reporting" guidance is now under development in OPP, and will specify how the study objectives, protocol, observations, findings, and conclusions are organized and presented within the study report. The data reporting guidance will be compatible with submittal format requirements described in this Notice.

OPP has also promulgated a policy (PR Notice 86-4 dated April 15, 1986) that provides for early screening of certain applications for registration under FIFRA §3. The objective of the screen is to avoid the additional costs and prolonged delays associated with handling significantly incomplete application packages. As of the effective date of this Notice, the screen will include in its criteria for acceptance of application packages the data formatting requirements described herein.

OPP has also established a public docket which imposes deadlines for inserting into the docket documents submitted in connection with Special Reviews and Registration Standards (see 40 CFR §154.15 and §155.32). To meet these deadlines, OPP is requiring an additional copy of any data submitted to the docket. Please refer to Page 10 for more information about this requirement.

For several years, OPP has required that each application for registration or other action include a list of all applicable data requirements and an indication of how each is satisfied-the statement of the method of support for the application. Typically, many requirements are satisfied by reference to data previously submitted-either by the applicant or by another party. That requirement is not altered by this notice, which applies only to data <u>submitted</u> with an application.

## VI. Format Requirements

A more detailed discussion of these format requirements follows the index on the next page, and samples of some of the requirements are attached. Except for the language of the two alternative forms of the Statement of Data Confidentiality Claims (shown in Attachment 3) which cannot be altered, these samples are illustrative. As long as the required information is included and clearly identifiable, the form of the samples may be altered to reflect the submitter's preference.

#### - INDEX-

A.	Page Organization of the Submittal Package	Page 1
В.	Transmittal Document 4	1
c.	Individual Studies 4	
	C. 1 Special Considerations for Identifying Studies 5	
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	D. 4 Supplemental Statement of Data Confidentiality	
	Claims (other than those based on FIFRA §10(d)(1)) 8	1
	D. 5 Good Laboratory Practice Compliance Statement 9	1
E.	Reference to Previously Submitted Data	
F.	Physical Format Requirements & Number of Copies 9	
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## A. Organization of Submittal Package

A "submittal package" consists of all studies submitted at the same time for review in support of a single regulatory action, along with a transmittal document and other related administrative material (e.g. the method of support statement, EPA Forms 8570-1, 8570-4, 8570-20, etc.) as appropriate.

Data submitters must organize each submittal package as described in this Notice. The transmittal and any other administrative material must be grouped together in the first physical volume. Each study included in the submittal package must then be bound separately.

Submitters sometimes provide additional materials that are intended to clarify, emphasize, or otherwise comment to help Product Managers and reviewers better understand the submittal.

- If such materials relate to <u>one</u> study, they should be included as an appendix to that study.
- If such materials relate to <u>more than one</u> study (as for example a summary of all studies in a discipline) or to the submittal in general, they must be included in the submittal package as a separate study (with title page and statement of confidentiality claims).



#### B. Transmittal Document

The first item in each submittal package must be a transmittal document. This document identifies the submitter or all joint submitters; the regulatory action in support of which the package is being submitted--i.e., a registration application, petition, experimental use permit (EUP), §3(c)(2)(B) data call-in, §6(a)(2) submittal, or a special review; the transmittal date; and a list of all individual studies included in the package in the order of their appearance, showing (usually by Guideline reference number) the data requirement(s) addressed by each one. The EPA-assigned number for the regulatory action (e.g. the registration, EUP, or tolerance petition number) should be included in the transmittal document as well, if it is known to the submitter. See Attachment 1 for an example of an acceptable transmittal document.

The list of included studies in the transmittal of a data submittal package supporting a registration application should be subdivided by discipline, reflecting the order in which data requirements appear in 40 CFR 158.

The list of included studies in the transmittal of a data submittal package supporting a petition for tolerance or an application for an EUP should be subdivided into sections A, B, C,... of the petition or application, as defined in 40 CFR 180.7 and 158.125, (petitions) or Pesticide Assessment Guidelines, Subdivision I (EUPs) as appropriate.

When a submittal package supports a tolerance petition <u>and</u> an application for a registration or an EUP, list the petition studies first, then the balance of the studies. Within these two groups of studies follow the instructions above.

## C. <u>Individual Studies</u>

A study is the report of a single scientific investigation, including all supporting analyses required for logical completeness. A study should be identifiable and distinguishable by a conventional bibliographic citation including author, date, and title. Studies generally correspond in scope to a single Guideline requirement for supporting data, with some exceptions discussed in section C.1. Each study included in a submittal package must be bound as a separate entity. (See comments on binding studies on page 9.)

Each study must be consecutively paginated, beginning from the title page as page 1. The total number of pages in the complete study must be shown on the study title page. In addition (to ensure that inadvertently separated pages can be reassociated with the proper study during handling or review) use either of the following:

- Include the total number of pages in the complete study on each page (i.e., 1 of 250, 2 of 250, ...250 of 250).
- Include a company name or mark and study number on each page of the study, e g , Company Name-1986-23. Never reuse a study number for marking the pages of subsequent studies.

When a single study is extremely long, binding it in multiple volumes is permissible so long as the entire study is paginated in a single series, and each volume is plainly identified by the study title and its position in the multi-volume sequence.

#### C.1 Special Considerations for Identifying Studies

Some studies raise special problems in study identification, because they address Guidelines of broader than normal scope or for other reasons.

a. <u>Safety Studies</u>. Several Guidelines require testing for safety in more than one species. In these cases each species tested should be reported as a separate study, and bound separately.

Extensive supplemental reports of pathology reviews, feed analyses, historical control data, and the like are often associated with safety studies. Whenever possible these should be submitted with primary reports of the study, and bound with the primary study as appendices. When such supplemental reports are submitted independently of the primary report, take care to fully identify the primary report to which they pertain.

Batteries of acute toxicity tests, performed on the same end use product and covered by a single title page, may be bound together and reported as a single study.

b. <u>Product Chemistry Studies</u>. All product chemistry data within a submittal package submitted in support of an end-use product produced from registered manufacturing-use products should be bound as a single study under a single title page.

Product chemistry data submitted in support of a technical product, other manufacturing-use product, an experimental use permit, an import tolerance petition, or an end-use product produced from unregistered source ingredients, should be bound as a single study for each Guideline <u>series</u> (61, 62, and 63) for conventional pesticides, or for the equivalent subject range for biorational pesticides. The first of the three studies in a complete product chemistry submittal for a biochemical pesticide would cover Guidelines 151-10, 151-11, and 151-12; the second would cover Guidelines 151-13, 151-15, and 151-16; the third would cover Guideline 151-17. The first study for a microbial pesticide would cover Guidelines 151-20, 151-21, and 151-22; the second would cover Guidelines 151-23 and 151-25; the third would cover Guideline 151-26.

Note particularly that product chemistry studies are likely to contain Confidential Business Information as defined in FIFRA §10(d)(1)(A), (B), or (C), and if so must be handled as described in section D.3. of this notice.



c. <u>Residue Chemistry Studies</u>. Guidelines 171-4, 153-3, and 153-4 are extremely broad in scope; studies addressing residue chemistry requirements must thus be defined at a level below that of the Guideline code. The general principle, however, of limiting a study to the report of a single investigation still applies fully. Data should be treated as a single study and bound separately for each analytical method, each report of the nature of the residue in a single crop or animal species, and for each report of the magnitude of residues resulting from treatment of a single crop or from processing a single crop. When more than one commodity is derived from a single crop (such as beet tops and beet roots) residue data on all such commodities should be reported as a single study. When multiple field trials are associated with a single crop, all such trials should be reported as a single study.

## D. Organization of Each Study Volume

Each complete study must include all applicable elements in the list below, in the order indicated. (Also see Page 17.) Several of these elements are further explained in the following paragraphs. Entries in the column headed "example" cite the page number of this notice where the element is illustrated.

<u>Element</u>	When Required	<u>Example</u>
Study Title Page	Always	Page 12
Statement of Data Confidentiality Claims	One of the two alternative forms of this statement is always required	Page 13
Certification of Good Laboratory Practice	If study reports laboratory work subject to GLP requirements	Page 16
Flagging statements	For certain toxicology studie flagging requirements are fin	s (When alized.)
Body of Study	Always - with an English lang translation if required.	uage
Study Appendices	At submitter's option	
Cover Sheet to Confidential Attachment	If CBI is claimed under FIFRA §10(d)(1)(A), (B), or (C)	
CBI Attachment	If CBI is claimed under FIFRA \$10(d)(1)(A), (B), or (C)	
Supplemental Statement of Data Confidentiality Claims	Only if confidentiality is claimed on a basis other than FIFRA §10(d)(1)(A), (B), or (	

### D.1. Title Page

A title page is always required for each submitted study, published or unpublished. The title page must always be freely releasable to requestors; DO NOT INCLUDE CBI ON THE TITLE PAGE. An example of an acceptable title page is on page 12 of this notice. The following information must appear on the title page:

- a. <u>Study title</u>. The study title should be as descriptive as possible It must clearly identify the substance(s) tested and correspond to the name of the data requirement as it appears in the Guidelines.
- b. <u>Data requirement addressed</u>. Include on the title page the Guideline number(s) of the specific requirement(s) addressed by the study.
- c. <u>Author(s)</u>. Cite only individuals with primary intellectual responsibility for the content of the study. Identify them plainly as authors, to distinguish them from the performing laboratory, study sponsor, or other names that may also appear on the title page.
- d. Study Date. The title page must include a single date for the study. If parts of the study were performed at different times, use only the date of the latest element in the study.
- e. <u>Performing Laboratory Identification</u>. If the study reports work done by one or more laboratories, include on the title page the name and address of the performing laboratory or laboratories, and the laboratory's internal project number(s) for the work. Clearly distinguish the laboratory's project identifier from any other reference numbers provided by the study sponsor or submitter.
- f. <u>Supplemental Submissions</u>. If the study is a commentary on or supplement to another previously submitted study, or if it responds to EPA questions raised with respect to an earlier study, include on the title page elements a. through d. for the previously submitted study, along with the EPA Master Record Identifier (MRID) or Accession number of the earlier study if you know these numbers. (Supplements submitted in the same submittal package as the primary study should be appended to and bound with the primary study. Do not include supplements to more than one study under a single title page).
- g. <u>Facts of Publication</u>. If the study is a reprint of a published document, identity on the title page all relevant facts of publication, such as the journal title, volume, issue, inclusive page numbers, and publication date.



D.2. Statements of Data Confidentiality Claims Under FIFRA §10(d)(1).

Each submitted study must be accompanied by one of the two alternative forms of the statement of Data Confidentiality Claims specified in the proposed regulation in §158.33 (b) and (c) (See Attachment 3). These statements apply only to claims of data confidentiality based on FIFRA §10(d)(1)(A), (B), or (C). Use the appropriate alternative form of the statement either to assert a claim of §10(d)(1) data confidentiality (§158.33(b)) or to waive such a claim (§158.33(c)). In either case, the statement must be signed and dated, and must include the typed name and title of the official who signs it. Do not make CBI claims with respect to analytical methods associated with petitions for tolerances or emergency exemptions (see NOTE Pg 13).

#### D.3. Confidential Attachment

If the claim is made that a study includes confidential business information as defined by the criteria of FIFRA §10(D)(1)(A), (B), or (C) (as described in D.2. above) all such information must be excised from the body of the study and confined to a separate study-specific Confidential Attachment. Each passage of CBI so isolated must be identified by a reference number cited within the body of the study at the point from which the passage was excised (See Attachment 5).

The Confidential Attachment to a study must be identified by a cover sheet fully identifying the parent study, and must be clearly marked "Confidential Attachment." An appropriately annotated photocopy of the parent study title page may be used as this cover sheet. Paginate the Confidential Attachment separately from the body of the study, beginning with page 1 of X on the title page. Each passage confined to the Confidential Attachment must be associated with a specific cross reference to the page(s) in the main body of the study on which it is cited, and with a reference to the applicable passage(s) of FIFRA §10(d)(1) on which the confidentiality claim is based.

D.4. <u>Supplemental</u> Statement of Data Confidentiality Claims (See Attachment 4)

If you wish to make a claim of confidentiality for any portion of a submitted study other than described by FIFRA §10(d)(1)(A), (B), or (C), the following provisions apply:

- The specific information to which the claim applies must be clearly marked in the body of the study as subject to a claim of confidentiality.
- A Supplemental Statement of Data Confidentiality Claims must be submitted, identifying each passage claimed confidential and describing in detail the basis for the claim. A list of the points to address in such a statement is included in Attachment 4 on Pg 14.
- The Supplemental Statement of Data Confidentiality Claims must be signed and dated and must include the typed name and title of the official who signed it.

## D.5. Good Laboratory Practice Compliance Statement

This statement is required if the study contains laboratory work subject to GLP requirements specified in 40 CFR 160. Samples of these statements are shown in Attachment 6.

#### E. Reference to Previously Submitted Data

FOR ANOTHER PURPOSE unless EPA specifically requests it. A copy of the title page plus the MRID number (if known) is sufficient to allow us to retrieve the study immediately for review. This prevents duplicate entries in the Agency files, and saves you the cost of sending more copies of the study. References to previously submitted studies should not be included in the transmittal document, but should be incorporated into the statement of the method of support for the application.

#### F. Physical Format Requirements

All elements in the data submittal package must be on uniform 8 1/2 by 11 inch white paper, printed on one side only in black ink, with high contrast and good resolution. Bindings for individual studies must be secure, but easily removable to permit disassembly for microfilming. Check with EPA for special instructions before submitting data in any medium other than paper, such as film or magnetic media.

Please be particularly attentive to the following points:

- Do not include frayed or torn pages.
- Do not include carbon copies, or copies in other than black ink.
- Make sure that photocopies are clear, complete, and fully readable.
- Do not include oversize computer printouts or fold-out pages.
- Do not bind any documents with glue or binding tapes.
- Make sure that all pages of each study, including any attachments or appendices, are present and in correct sequence.

Number of Copies Required - All submittal packages except those associated with a Registration Standard or Special Review (See Part G below) must be provided in three complete, identical copies. (The proposed regulations specified two copies; three are now being required to expedite and reduce the cost of processing data into the OPP Pesticide Document Management System and getting it into review.)



## G. Special Requirements for Submitting Data to the Docket

Data submittal packages associated with a Registration Standard or Special Review must be provided in <u>four</u> copies, from one of which all material claimed as CBI has been excised. This fourth copy will become part of the public docket for the RS or SR case. If no claims of confidentiality are made for the study, the fourth copy should be identical to the other three. When portions of a study submitted in support of an RS or SR are claimed as CBI, the first three copies will include the CBI material as provided in section D of this notice. The following special preparation is required for the fourth copy.

- Remove the "Supplemental Statement of Data Confidentiality Claims".
- Remove the "Confidential Attachment".
- Excise from the body of the study any information you claim as confidential, even if it does not fall within the scope of FIFRA §10(d)(1)(A), (B), or (C). Do not close up or paraphrase text remaining after this excision.
- Mark the fourth copy plainly on both its cover and its title page with the phrase "Public Docket Material contains no information claimed as confidential".

### V. <u>For Further Information</u>

For further information contact John Carley, Chief, Information Services Branch, Program Management and Support Division, (703) 305-5240.

Attachment 1. Sample Transmittal Document Attachment 2. Sample Title Page for a Newly Submitted Study

Tames W. Akerman Acting Director.

Registration Division

Attachment 3. Statements of Data Confidentiality Claims

Attachment 4. Supplemental Statement of Data Confidentiality Claims

Attachment 5. Samples of Confidential Attachments

Attachment 6. Sample Good Laboratory Practice Statements

Attachment 7. Format Diagrams for Submittal Packages and Studies

## ELEMENTS TO BE INCLUDED IN THE TRANSMITTAL DOCUMENT\*

1. Name and address of submitter (or all joint submitters\*\*)

\*Smith Chemical Corporation Jones Chemical Company 1234 West Smith Street -and-Cincinnati, OH 98765 Jones Chemical Company 5678 Wilson Blvd Covington, KY 56789

\*Smith Chemical Corp will act as sole agent for all submitters.

2. Regulatory action in support of which this package is submitted

Use the EPA identification number (e.g. 359-EUP-67) if you know it. Otherwise describe the type of request (e.g. experimental use permit, data call-in - of xx-xx-xx date).

- 3. Transmittal date
- 4. List of submitted studies
  - Vol 1. Administrative materials forms, previous correspondence with Project Managers, and so forth.
  - Vol 2. Title of first study in the submittal (Guideline No.)
  - Vol n Title of nth study in the submittal (Guideline No.)
  - \* Applicants commonly provide this information in a transmittal letter. This remains an acceptable practice so long as all four elements are included.
  - \* Indicate which of the joint submitters is empowered to act on behalf of all joint submitters in any matter concerning data compensation or subsequent use or release of the data.

Company	Official:		
		Name	Signature
Company	Name:		
Company .	Contact:	•	
		Name	Phone



#### SAMPLE STUDY TITLE PAGE FOR A NEWLY SUBMITTED STUDY

## Study Title

(Chemical name) - Magnitude of Residue on Corn

Data Requirement

Guideline 171-4

Author

John C. Davis

Study Completed On

January 5, 1979

Performing Laboratory

ABC Agricultural Laboratories 940 West Bay Drive Wilmington, CA 39897

Laboratory Project ID

ABC 47-79

Page 1 of X (X is the total number of pages in the study)

#### STATEMENTS OF DATA CONFIDENTIALITY CLAIMS

No claim of confidentiality under FIFRA §10(d)(1)(A),(B), or
 (C).

#### STATEMENT OF NO DATA CONFIDENTIALITY CLAIMS

	dy on the basis of	its falling within
Company		
STATEMENT OF DATA CONFIDENTIALITY CLAIMS  Information claimed confidential on the basis of its falling within the scope of FIFRA §10(d)(1)(A), (B), or (C) has been		
Title		Signature
(C).		
Information claimed c within the scope of F removed to a confiden cross-reference numbe	TFRA §10(d)(1)(A), tial appendix, and	(B), or (C) has been is cited by
Company:		
Company Agent:	Typed Name	Date:
Title		Signature

NOTE: Applicants for permanent or temporary tolerances should note that it is OPP policy that no permanent tolerance, temporary tolerance, or request for an emergency exemption incorporating an analytical method, can be approved unless the applicant waives all claims of confidentiality for the analytical method. These analytical methods are published in the FDA Pesticide Analytical Methods Manual, and therefore cannot be claimed as confidential. OPP implements this policy by returning submitted analytical methods, for which confidentiality claims have been made, to the submitter, to obtain the confidentiality waiver before they can be processed.



#### SUPPLEMENTAL STATEMENT OF DATA CONFIDENTIALITY CLAIMS

For any portion of a submitted study that is not described by FIFRA  $\S10(d)(1)(A)$ , (B), or (C), but for which you claim confidential treatment on another basis, the following information must be included within a Supplemental Statement of Data Confidentiality Claims:

- Identify specifically by page and line number(s) each portion of the study for which you claim confidentiality.
- Cite the reasons why the cited passage qualifies for confidential treatment.
- Indicate the length of time--until a specific date or event, or permanently--for which the information should be treated as confidential.
- Identify the measures taken to guard against undesired disclosure of this information.
- Describe the extent to which the information has been disclosed, and what precautions have been taken in connection with those disclosures.
- Enclose copies of any pertinent determinations of confidentiality made by EPA, other Federal agencies, of courts concerning this information.
- If you assert that disclosure of this information would be likely to result in substantial harmful effects to you, describe those harmful effects and explain why they should be viewed as substantial.
- If you assert that the information in voluntarily submitted, indicate whether you believe disclosure of this information might tend to lessen the availability to EPA of similar information in the future, and if so, how.

## EXAMPLES OF SEVERAL CONFIDENTIAL ATTACHMENTS

Example 1. (Confidential word or phrase that has been deleted from the study)

CROSS F	REFERENC	E NUMBER 1	in place	of the fol:	e number is lowing words d page refer	or ph	
DELETE	WORDS	OR PHRASE:		Ethyler	ne Glycol		<del></del>
PAGE	LINE	REASON FOR	THE DELE	TION		FIFRA	REFERENCE
6	14	Identity of	Inert I	ngredient		§10 (d)	(1) (C)
12	25	-	11	*	* •		н
100	19		Ħ		•		И

## Example 2. (Confidential paragraph(s) that have been deleted from the study)

CROSS REF	TERENCE NUMBER 5 This cross reference number is used in the in place of the following paragraph(s) are indicated volume and page references.	
DELETED	PARAGRAPH(S):	
(		)
(	Reproduce the deleted paragraph(s) here	)
(	•	)
PAGE	LINES REASON FOR THE DELETION FIFRA R	EFERENCE
20.	2-17 Description of the quality control process \$10(	d) (1) (C)

#### Example 3. (Confidential pages that have been deleted from the study)

CROSS REFERENCE NUMBER 7 This cross reference number noted on a placeholder page is used in place of the following
whole pages at the indicated volume and page
references.

DELETED PAGE(S): are attached immediately behind this page.

PAGE LINES REASON FOR THE DELETION FIFRA REFERENCE

20. 2-17 Description of the product manufacturing process \$10 (d) (1) (A)



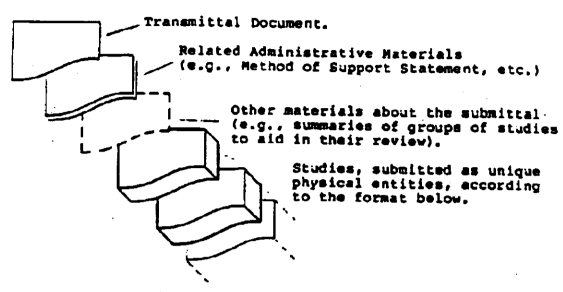
## ATTACHMENT 6.

## SAMPLE GOOD LABORATORY PRACTICE STATEMENTS

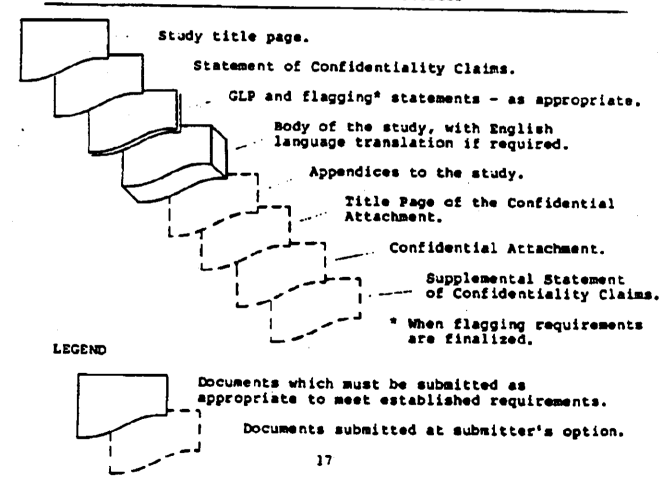
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#### ATTACHMENT 7.

## FORMAT OF THE SUBMITTAL PACKAGE



## FORMAT OF SUBMITTED STUDIES



PR Notice 91-2



## UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

WASHINGTON, D.C. 20460

OFFICE OF
PREVENTION, PESTICIDES
AND TOXIC SUBSTANCES

#### PR NOTICE 91-2

NOTICE TO MANUFACTURERS, PRODUCERS, FORMULATORS, AND REGISTRANTS OF PESTICIDES

ATTENTION: Persons Responsible for Federal Registration of Pesticide Products.

SUBJECT: Accuracy of Stated Percentages for Ingredients Statement

#### I. PURPOSE:

The purpose of this notice is to clarify the Office of Pesticide Program's policy with respect to the statement of percentages in a pesticide's label's ingredient statement. Specifically, the amount (percent by weight) of ingredient(s) specified in the ingredient statement on the label must be stated as the nominal concentration of such ingredient(s), as that term is defined in 40 CFR 158.153(i). Accordingly, the Agency has established the nominal concentration as the only acceptable label claim for the amount of active ingredient in the product.

#### II. BACKGROUND

For some time the Agency has accepted two different methods of identifying on the label what percentage is claimed for the ingredient(s) contained in a pesticide. Some applicants claimed a percentage which represented a level between the upper and the lower certified limits. This was referred to as the nominal concentration. Other applicants claimed the lower limit as the percentage of the ingredient(s) that would be expected to be present in their product at the end of the product's shelf-life. Unfortunately, this led to a great deal of confusion among the regulated industry, the regulators, and the consumers as to exactly how much of a given ingredient was in a given product. The Agency has established the nominal concentration as the only acceptable label claim for the amount of active ingredient in the product.

Current regulations require that the percentage listed in the active ingredient statement be as precise as possible reflecting good manufacturing practices 40 CFR 156.10(g)(5). The



certified limits required for each active ingredient are intended to encompass any such "good manufacturing practice" variations 40 CFR 158.175(c)(3).

The upper and lower certified limits, which must be proposed in connection with a product's registration, represent the amounts of an ingredient that may legally be present 40 CFR 158.175. The lower certified limit is used as the enforceable lower limit for the product composition according to FIFRA section 12(a)(1)(C), while the nominal concentration appearing on the label would be the routinely achieved concentration used for calculation of dosages and dilutions.

The nominal concentration would in fact state the greatest degree of accuracy that is warranted with respect to actual product composition because the nominal concentration would be the amount of active ingredient typically found in the product.

It is important for registrants to note that certified limits for active ingredients are not considered to be trade secret information under FIFRA section 10(b). In this respect the certified limits will be routinely provided by EPA to States for enforcement purposes, since the nominal concentration appearing on the label may not represent the enforceable composition for purposes of section 12(a)(1)(C).

#### III. REQUIREMENTS

As described below under Unit V. " COMPLIANCE SCHEDULE, " all currently registered products as well as all applications for new registration must comply with this Notice by specifying the nominal concentration expressed as a percentage by weight as the label claim in the ingredient(s) statement and equivalence statements if applicable (e.g., elemental arsenic, metallic zinc, salt of an acid). In addition, the requirement for performing sample analyses of five or more representative samples must be fulfilled. Copies of the raw analytical data must be submitted with the nominal ingredient label claim. Further information about the analysis requirement may be found in the 40 CFR 158.170. All products are required to provide certified limits for each active, inert ingredient, impurities of toxicological significance(i.e., upper limit(s) only) and on a case by case basis as specified by EPA. These limits are to be set based on representative sampling and chemical analysis (i.e., guality control) of the product.

The format of the ingredient statement must conform to 40 CFR 156-Labeling Requirements For Pesticides and Devices.

After July 1, 1997, all pesticide ingredient StatementS must be changed to nominal concentration.

### IV. PRODUCTS THAT REQUIRE EFFICACY DATA

All pesticides are required to be efficacious. Therefore, the certified lower limits may not be lower then the minimum level to achieve efficacy. This is extremely important for products which are intended to control pests which threaten the public health, e.g., certain antimicrobial and rodenticide products. Refer to 40 CFR 153.640.

In those cases where efficacy limits have been established, the Agency will not accept certified lower limits which are below that level for the shelf life of the product.

#### V. COMPLIANCE SCHEDULE

As described earlier, the purpose of this Notice is to make the registration process more uniform and more manageable for both the agency and the regulated community. It is the Agency's intention to implement the requirements of this notice as smoothly as possible so as not to disrupt or delay the Agency's high priority programs, i.e., reregistration, new chemical, or fast track (FIFRA section 3(c)(3)(B). Therefore, applicants/registrants are expected to comply with the requirements of this Notice as follows:

- (1) Beginning July 1, 1991, all new product registrations submitted to the Agency are to comply with the requirements of this Notice.
- (2) Registrants having products subject to reregistration under FIFRA section 4(a) are to comply with the requirements of this Notice when specific products are called in by the Agency under Phase V of the Reregistration Program.
- (3) All other products/applications that are not subject to (1) and (2) above will have until July 1, 1997, to comply with this Notice. Such applications should note "Conversion to Nominal Concentrations on the application form. These types Or amendments will not be handled as "Fast Track" applications but will be handled as routine requests.

#### VI. FOR FURTHER INFORMATION

Contact Tyrone Aiken for information or questions concerning this notice on (703) 308-7031.

Anne E. Lindsay, Director Registration Division (H-7505 

# APPENDIX F Generic Data Call-In

#### DATA CALL-IN NOTICE

#### **CERTIFIED MAIL**

#### Dear Sir or Madam:

This Notice requires you and other registrants of pesticide products containing the active ingredient(s) identified in Attachment A of this Notice, the <u>Data Call-In Chemical Status Sheet</u>, to submit certain data as noted herein to the U.S. Environmental Protection Agency (EPA, the Agency). These data are necessary to maintain the continued registration of your product(s) containing this active ingredient(s). Within 90 days after you receive this Notice you must respond as set forth in Section III below. Your response must state:

- 1. how you will comply with the requirements set forth in this Notice and its Attachments A through D; or,
- 2. why you believe you are exempt from the requirements listed in this Notice and in Attachment C, Requirements Status and Registrant's Response Form, (see section III-B); or.
- 3. why you believe EPA should not require your submission of data in the manner specified by this Notice (see section III-D).

If you do not respond to this Notice, or if you do not satisfy EPA that you will comply with its requirements or should be exempt or excused from doing so, then the registration of your product(s) subject to this Notice will be subject to suspension. We have provided a list of all of your products subject to this Notice in Attachment B, <u>Data Call-In Response Form</u>, as well as a list of all registrants who were sent this Notice (Attachment D).

The authority for this Notice is section 3(c)(2)(B) of the Federal Insecticide, Fungicide and Rodenticide Act as amended (FIFRA), 7 U.S.C. section 136a(c)(2)(B). Collection of this information is authorized under the Paperwork Reduction Act by OMB Approval No. 2070-0107 (expiration date 12-31-92).



This Notice is divided into six sections and five Attachments. The Notice itself contains information and instructions applicable to all Data Call-In Notices. The Attachments contain specific chemical information and instructions. The six sections of the Notice are:

Section I - Why You Are Receiving This Notice

Section II - Data Required By This Notice

Section III - Compliance With Requirements Of This Notice

Section IV - Consequences Of Failure To Comply With This Notice

Section V - Registrants' Obligation To Report Possible Unreasonable Adverse

**Effects** 

Section VI - Inquiries And Responses To This Notice

# The Attachments to this Notice are:

Attachment A - Data Call-In Chemical Status Sheet

Attachment B - Data Call-In Response Form

Attachment C - Requirements Status And Registrant's Response Form

Attachment D - List Of All Registrants Sent This Data Call-In Notice

# SECTION I. WHY YOU ARE RECEIVING THIS NOTICE

The Agency has reviewed existing data for this active ingredient(s) and reevaluated the data needed to support continued registration of the subject active ingredient(s). This reevaluation identified additional data necessary to assess the health and safety of the continued use of products containing this active ingredient(s). You have been sent this Notice because you have product(s) containing the subject active ingredient(s).

### SECTION II. DATA REQUIRED BY THIS NOTICE

# A. <u>DATA REQUIRED</u>

The data required by this Notice are specified in Attachment C, <u>Requirements Status and Registrant's Response Form</u>. Depending on the results of the studies required in this Notice, additional testing may be required.

# B. SCHEDULE FOR SUBMISSION OF DATA

You are required to submit the data or otherwise satisfy the data requirements specified in Attachment C, Requirements Status and Registrant's Response Form, within the time frames provided.

# C. TESTING PROTOCOL

All studies required under this Notice must be conducted in accordance with test standards outlined in the Pesticide Assessment Guidelines for those studies for which guidelines have been established.

These EPA Guidelines are available from the National Technical Information Service (NTIS), Attn: Order Desk, 5285 Port Royal Road, Springfield, Va 22161 (tel: 703-487-4650).

Protocols approved by the Organization for Economic Cooperation and Development (OECD) are also acceptable if the OECD-recommended test standards conform to those specified in the Pesticide Data Requirements regulation (40 CFR § 158.70). When using the OECD protocols, they should be modified as appropriate so that the data generated by the study will satisfy the requirements of 40 CFR § 158. Normally, the Agency will not extend deadlines for complying with data requirements when the studies were not conducted in accordance with acceptable standards. The OECD protocols are available from OECD, 1750 Pennsylvania Avenue N.W., Washington, D.C. 20006.

All new studies and proposed protocols submitted in response to this Data Call-In Notice must be in accordance with Good Laboratory Practices [40 CFR Part 160.3(a)(6)].

# D. <u>REGISTRANTS RECEIVING PREVIOUS SECTION 3(c)(2)(B) NOTICES</u> ISSUED BY THE AGENCY

Unless otherwise noted herein, this Data Call-In does not in any way supersede or change the requirements of any previous Data Call-In(s), or any other agreements entered into with the Agency pertaining to such prior Notice. Registrants must comply with the requirements of all Notices to avoid issuance of a Notice of Intent to Suspend their affected products.

# SECTION III. COMPLIANCE WITH REQUIREMENTS OF THIS NOTICE

# A. SCHEDULE FOR RESPONDING TO THE AGENCY

The appropriate responses initially required by this Notice must be submitted to the Agency within 90 days after your receipt of this Notice. Failure to adequately respond to this Notice within 90 days of your receipt will be a basis for issuing a Notice of Intent to Suspend (NOIS) affecting your products. This and other bases for issuance of NOIS due to failure to comply with this Notice are presented in Section IV-A and IV-B.



# B. OPTIONS FOR RESPONDING TO THE AGENCY

The options for responding to this Notice are: 1) voluntary cancellation, 2) delete use(s), (3) claim generic data exemption, (4) agree to satisfy the data requirements imposed by this Notice or (5) request a data waiver(s).

A discussion of how to respond if you chose the Voluntary Cancellation option, the Delete Use(s) option or the Generic Data Exemption option is presented below. A discussion of the various options available for satisfying the data requirements of this Notice is contained in Section III-C. A discussion of options relating to requests for data waivers is contained in Section III-D.

There are two forms that accompany this Notice of which, depending upon your response, one or both must be used in your response to the Agency. These forms are the <u>Data-Call-In Response Form</u> (Attachment B) and the <u>Requirements Status and Registrant's Response Form</u> (Attachment C). The <u>Data Call-In Response Form</u> must be submitted as part of every response to this Notice. Please note that the company's authorized representative is required to sign the first page of the <u>Data Call-In Response Form</u> and <u>Requirements Status and Registrant's Response Form</u> (if this form is required) and initial any subsequent pages. The forms contain separate detailed instructions on the response options. Do not alter the printed material. If you have questions or need assistance in preparing your response, call or write the contact person identified in Attachment A.

1. <u>Voluntary Cancellation</u> - You may avoid the requirements of this Notice by requesting voluntary cancellation of your product(s) containing the active ingredient(s) that is the subject of this Notice. If you wish to voluntarily cancel your product, you must submit a completed <u>Data Call-In Response Form</u>, indicating your election of this option. Voluntary cancellation is item number 5 on the <u>Data Call-In Response Form</u>. If you choose this option, this is the only form that you are required to complete.

If you choose to voluntarily cancel your product, further sale and distribution of your product after the effective date of cancellation must be in accordance with the Existing Stocks provisions of this Notice which are contained in Section IV-C.

2. <u>Use Deletion</u> - You may avoid the requirements of this Notice by eliminating the uses of your product to which the requirements apply. If you wish to amend your registration to delete uses, you must submit the <u>Requirements Status and Registrant's Response Form</u>, a completed application for amendment, a copy of your proposed amended labeling, and all other information required for processing the application. Use deletion is option number 7 on the <u>Requirements Status and Registrant's Response Form</u>. You must also complete a <u>Data Call-In</u>

Response Form by signing the certification, item number 8. Application forms for amending registrations may be obtained from the Registration Support and Emergency Response Branch, Registration Division, (703) 308-8358.

If you choose to delete the use(s) subject to this Notice or uses subject to specific data requirements, further sale, distribution, or use of your product after one year from the due date of your 90 day response, must bear an amended label.

- 3. Generic Data Exemption Under section 3(c)(2)(D) of FIFRA, an applicant for registration of a product is exempt from the requirement to submit or cite generic data concerning an active ingredient(s) if the active ingredient(s) in the product is derived exclusively from purchased, registered pesticide products containing the active ingredient(s). EPA has concluded, as an exercise of its discretion, that it normally will not suspend the registration of a product which would qualify and continue to qualify for the generic data exemption in section 3(c)(2)(D) of FIFRA. To qualify, <u>all</u> of the following requirements must be met:
  - a. The active ingredient(s) in your registered product must be present solely because of incorporation of another registered product which contains the subject active ingredient(s) and is purchased from a source not connected with you; and,
  - b. every registrant who is the ultimate source of the active ingredient(s) in your product subject to this DCI must be in compliance with the requirements of this Notice and must remain in compliance; and
  - c. you must have provided to EPA an accurate and current "Confidential Statement of Formula" for each of your products to which this Notice applies.

To apply for the Generic Data Exemption you must submit a completed <u>Data Call-In Response Form</u>, Attachment B and all supporting documentation. The Generic Data Exemption is item number 6a on the <u>Data Call-In Response Form</u>. If you claim a generic data exemption you are not required to complete the <u>Requirements Status and Registrant's Response Form</u>. Generic Data Exemption cannot be selected as an option for product specific data.

If you are granted a Generic Data Exemption, you rely on the efforts of other persons to provide the Agency with the required data. If the registrant(s) who have committed to generate and submit the required data fail to take appropriate steps to meet the requirements or are no longer in compliance with this Data Call-In Notice, the Agency will consider that both they and you are not in compliance and will normally initiate proceedings to suspend the registrations



of both your and their product(s), unless you commit to submit and do submit the required data within the specified time. In such cases the Agency generally will not grant a time extension for submitting the data.

- 4. <u>Satisfying the Data Requirements of this Notice</u> There are various options available to satisfy the data requirements of this Notice. These options are discussed in Section III-C of this Notice and comprise options 1 through 6 on the <u>Requirements Status and Registrant's Response Form</u> and option 6b and 7 on the <u>Data Call-In Response Form</u>. If you choose option 6b or 7, you must submit both forms as well as any other information/data pertaining to the option chosen to address the data requirement.
- 5. Request for Data Waivers. Data waivers are discussed in Section III-D of this Notice and are covered by options 8 and 9 on the Requirements Status and Registrant's Response Form. If you choose one of these options, you must submit both forms as well as any other information/data pertaining to the option chosen to address the data requirement.

# C. SATISFYING THE DATA REQUIREMENTS OF THIS NOTICE

If you acknowledge on the <u>Data Call-In Response Form</u> that you agree to satisfy the data requirements (i.e. you select option 6b and/or 7), then you must select one of the six options on the <u>Requirements Status and Registrant's Response Form</u> related to data production for each data requirement. Your option selection should be entered under item number 9, "Registrant Response." The six options related to data production are the first six options discussed under item 9 in the instructions for completing the <u>Requirements Status and Registrant's Response Form</u>. These six options are listed immediately below with information in parentheses to guide registrants to additional instructions provided in this Section. The options are:

- 1. I will generate and submit data within the specified time frame (Developing Data),
- 2. I have entered into an agreement with one or more registrants to develop data jointly (Cost Sharing),
- 3. I have made offers to cost-share (Offers to Cost Share),
- 4. I am submitting an existing study that has not been submitted previously to the Agency by anyone (Submitting an Existing Study),
- 5. I am submitting or citing data to upgrade a study classified by EPA as partially acceptable and upgradeable (Upgrading a Study),

6. I am citing an existing study that EPA has classified as acceptable or an existing study that has been submitted but not reviewed by the Agency (Citing an Existing Study).

# Option 1, Developing Data --

If you choose to develop the required data it must be in conformance with Agency deadlines and with other Agency requirements as referenced herein and in the attachments. All data generated and submitted must comply with the Good Laboratory Practice (GLP) rule (40 CFR Part 160), be conducted according to the Pesticide Assessment Guidelines (PAG), and be in conformance with the requirements of PR Notice 86-5. In addition, certain studies require Agency approval of test protocols in advance of study initiation. Those studies for which a protocol must be submitted have been identified in the Requirements Status and Registrant's Response Form and/or footnotes to the form. If you wish to use a protocol which differs from the options discussed in Section II-C of this Notice, you must submit a detailed description of the proposed protocol and your reason for wishing to use it. The Agency may choose to reject a protocol not specified in Section II-C. If the Agency rejects your protocol you will be notified in writing, however, you should be aware that rejection of a proposed protocol will not be a basis for extending the deadline for submission of data.

A progress report must be submitted for each study within 90 days from the date you are required to commit to generate or undertake some other means to address that study requirement, such as making an offer to cost-share or agreeing to share in the cost of developing that study. A 90-day progress report must be submitted for all studies. This 90-day progress report must include the date the study was or will be initiated and, for studies to be started within 12 months of commitment, the name and address of the laboratory(ies) or individuals who are or will be conducting the study.

In addition, if the time frame for submission of a final report is more than 1 year, interim reports must be submitted at 12 month intervals from the date you are required to commit to generate or otherwise address the requirement for the study. In addition to the other information specified in the preceding paragraph, at a minimum, a brief description of current activity on and the status of the study must be included as well as a full description of any problems encountered since the last progress report.

The time frames in the Requirements Status and Registrant's Response Form are the time frames that the Agency is allowing for the submission of completed study reports or protocols. The noted deadlines run from the date of the receipt of this Notice by the registrant. If the data are not submitted by the deadline, each registrant is subject to receipt of a Notice of Intent to Suspend the

# affected registration(s).

If you cannot submit the data/reports to the Agency in the time required by this Notice and intend to seek additional time to meet the requirement(s), you must submit a request to the Agency which includes: (1) a detailed description of the expected difficulty and (2) a proposed schedule including alternative dates for meeting such requirements on a step-by-step basis. You must explain any technical or laboratory difficulties and provide documentation from the laboratory performing the testing. While EPA is considering your request, the original deadline remains. The Agency will respond to your request in writing. If EPA does not grant your request, the original deadline remains. Normally, extensions can be requested only in cases of extraordinary testing problems beyond the expectation or control of the registrant. Extensions will not be given in submitting the 90-day responses. Extensions will not be considered if the request for extension is not made in a timely fashion; in no event shall an extension request be considered if it is submitted at or after the lapse of the subject deadline.

# Option 2, Agreement to Share in Cost to Develop Data --

If you choose to enter into an agreement to share in the cost of producing the required data but will not be submitting the data yourself, you must provide the name of the registrant who will be submitting the data. You must also provide EPA with documentary evidence that an agreement has been formed. Such evidence may be your letter offering to join in an agreement and the other registrant's acceptance of your offer, or a written statement by the parties that an agreement exists. The agreement to produce the data need not specify all of the terms of the final arrangement between the parties or the mechanism to resolve the terms. Section 3(c)(2)(B) provides that if the parties cannot resolve the terms of the agreement they may resolve their differences through binding arbitration.

# Option 3, Offer to Share in the Cost of Data Development --

If you have made an offer to pay in an attempt to enter into an agreement or amend an existing agreement to meet the requirements of this Notice and have been unsuccessful, you may request EPA (by selecting this option) to exercise its discretion not to suspend your registration(s), although you do not comply with the data submission requirements of this Notice. EPA has determined that as a general policy, absent other relevant considerations, it will not suspend the registration of a product of a registrant who has in good faith sought and continues to seek to enter into a joint data development/cost sharing program, but the other registrant(s) developing the data has refused to accept your offer. To qualify for this option, you must submit documentation to the Agency proving that you have made an offer to another registrant (who has an obligation to submit



data) to share in the burden of developing that data. You must also submit to the Agency a completed EPA Form 8570-32, Certification of Offer to Cost Share in the Development of Data, Attachment E. In addition, you must demonstrate that the other registrant to whom the offer was made has not accepted your offer to enter into a cost sharing agreement by including a copy of your offer and proof of the other registrant's receipt of that offer (such as a certified mail receipt). Your offer must, in addition to anything else, offer to share in the burden of producing the data upon terms to be agreed or failing agreement to be bound by binding arbitration as provided by FIFRA section 3(c)(2)(B)(iii) and must not qualify this offer. The other registrant must also inform EPA of its election of an option to develop and submit the data required by this Notice by submitting a Data Call-In Response Form and a Requirements Status and Registrant's Response Form committing to develop and submit the data required by this Notice.

In order for you to avoid suspension under this option, you may not withdraw your offer to share in the burdens of developing the data. In addition, the other registrant must fulfill its commitment to develop and submit the data as required by this Notice. If the other registrant fails to develop the data or for some other reason is subject to suspension, your registration as well as that of the other registrant will normally be subject to initiation of suspension proceedings, unless you commit to submit, and do submit the required data in the specified time frame. In such cases, the Agency generally will not grant a time extension for submitting the data.

# Option 4, Submitting an Existing Study --

If you choose to submit an existing study in response to this Notice, you must determine that the study satisfies the requirements imposed by this Notice. You may only submit a study that has not been previously submitted to the Agency or previously cited by anyone. Existing studies are studies which predate issuance of this Notice. Do not use this option if you are submitting data to upgrade a study. (See Option 5).

You should be aware that if the Agency determines that the study is not acceptable, the Agency will require you to comply with this Notice, normally without an extension of the required date of submission. The Agency may determine at any time that a study is not valid and needs to be repeated.

To meet the requirements of the DCI Notice for submitting an existing study, all of the following three criteria must be clearly met:

- You must certify at the time that the existing study is submitted a. that the raw data and specimens from the study are available for audit and review and you must identify where they are available. This must be done in accordance with the requirements of the Good Laboratory Practice (GLP) regulation, 40 CFR Part 160. As stated in 40 CFR 160.3(7) " raw data means any laboratory worksheets, records, memoranda, notes, or exact copies thereof, that are the result of original observations and activities of a study and are necessary for the reconstruction and evaluation of the report of that study. In the event that exact transcripts of raw data have been prepared (e.g., tapes which have been transcribed verbatim, dated, and verified accurate by signature), the exact copy or exact transcript may be substituted for the original source as raw data. Raw data may include photographs, microfilm or microfiche copies, computer printouts, magnetic media, including dictated observations, and recorded data from automated instruments." The term "specimens". according to 40 CFR 160.3(7), means "any material derived from a test system for examination or analysis."
- b. Health and safety studies completed after May 1984 must also contain all GLP-required quality assurance and quality control information, pursuant to the requirements of 40 CFR Part 160. Registrants must also certify at the time of submitting the existing study that such GLP information is available for post-May 1984 studies by including an appropriate statement on or attached to the study signed by an authorized official or representative of the registrant.
- You must certify that each study fulfills the acceptance criteria for c. the Guideline relevant to the study provided in the FIFRA Accelerated Reregistration Phase 3 Technical Guidance and that the study has been conducted according to the Pesticide Assessment Guidelines (PAG) or meets the purpose of the PAG (both available from NTIS). A study not conducted according to the PAG may be submitted to the Agency for consideration if the registrant believes that the study clearly meets the purpose of the PAG. The registrant is referred to 40 CFR 158.70 which states the Agency's policy regarding acceptable protocols. If you wish to submit the study, you must, in addition to certifying that the purposes of the PAG are met by the study, clearly articulate the rationale why you believe the study meets the purpose of the PAG, including copies of any supporting information or data. It has been the Agency's experience that studies-completed prior to January 1970 rarely satisfied the purpose of the PAG and that necessary raw data are usually not available for such studies.

If you submit an existing study, you must certify that the study meets all requirements of the criteria outlined above.

If EPA has previously reviewed a protocol for a study you are submitting, you must identify any action taken by the Agency on the protocol and must indicate, as part of your certification, the manner in which all Agency comments, concerns, or issues were addressed in the final protocol and study.

If you know of a study pertaining to any requirement in this Notice which does not meet the criteria outlined above but does contain factual information regarding unreasonable adverse effects, you must notify the Agency of such a study. If such a study is in the Agency's files, you need only cite it along with the notification. If not in the Agency's files, you must submit a summary and copies as required by PR Notice 86-5.

# Option 5, Upgrading a Study --

If a study has been classified as partially acceptable and upgradeable, you The Agency will review the data may submit data to upgrade that study. submitted and determine if the requirement is satisfied. If the Agency decides the requirement is not satisfied, you may still be required to submit new data normally without any time extension. Deficient, but upgradeable studies will normally be classified as supplemental. However, it is important to note that not all studies classified as supplemental are upgradeable. If you have questions regarding the classification of a study or whether a study may be upgraded, call or write the contact person listed in Attachment A. If you submit data to upgrade an existing study you must satisfy or supply information to correct all deficiencies in the study identified by EPA. You must provide a clearly articulated rationale of how the deficiencies have been remedied or corrected and why the study should be rated as acceptable to EPA. Your submission must also specify the MRID number(s) of the study which you are attempting to upgrade and must be in conformance with PR Notice 86-5.

Do not submit additional data for the purpose of upgrading a study classified as unacceptable and determined by the Agency as not capable of being upgraded.

This option should also be used to cite data that has been previously submitted to upgrade a study, but has not yet been reviewed by the Agency. You must provide the MRID number of the data submission as well as the MRID number of the study being upgraded.

The criteria for submitting an existing study, as specified in Option 4

above, apply to all data submissions intended to upgrade studies. Additionally your submission of data intended to upgrade studies must be accompanied by a certification that you comply with each of those criteria as well as a certification regarding protocol compliance with Agency requirements.

# Option 6, Citing Existing Studies --

If you choose to cite a study that has been previously submitted to EPA, that study must have been previously classified by EPA as acceptable or it must be a study which has not yet been reviewed by the Agency. Acceptable toxicology studies generally will have been classified as "core-guideline" or "core minimum." For ecological effects studies, the classification generally would be a rating of "core." For all other disciplines the classification would be "acceptable." With respect to any studies for which you wish to select this option you must provide the MRID number of the study you are citing and, if the study has been reviewed by the Agency, you must provide the Agency's classification of the study.

If you are citing a study of which you are not the original data submitter, you must submit a completed copy of EPA Form 8570-31, Certification with Respect to Data Compensation Requirements.

# D. REQUESTS FOR DATA WAIVERS

There are two types of data waiver responses to this Notice. The first is a request for a low volume/minor use waiver and the second is a waiver request based on your belief that the data requirement(s) are inapplicable and do not apply to your product.

Low Volume/Minor Use Waiver -- Option 8 on the Requirements Status and Registrant's Response Form. Section 3(c)(2)(A) of FIFRA requires EPA to consider the appropriateness of requiring data for low volume, minor use In implementing this provision EPA considers as low volume pesticides only those active ingredient(s) whose total production volume for all pesticide registrants is small. In determining whether to grant a low volume, minor use waiver the Agency will consider the extent, pattern and volume of use. the economic incentive to conduct the testing, the importance of the pesticide, and the exposure and risk from use of the pesticide. If an active ingredient(s) is used for both high volume and low volume uses, a low volume exemption will not be approved. If all uses of an active ingredient(s) are low volume and the combined volumes for all-uses are also low, then an exemption may be granted, depending on review of other information outlined below. An exemption will not be granted if any registrant of the active ingredient(s) elects to conduct the testing. Any registrant receiving a low volume minor use waiver must remain within the sales figures in their forecast supporting the waiver request in order to remain qualified



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for such waiver. If granted a waiver, a registrant will be required, as a condition of the waiver, to submit annual sales reports. The Agency will respond to requests for waivers in writing.

To apply for a low volume, minor use waiver, you must submit the following information, as applicable to your product(s), as part of your 90-day response to this Notice:

- a. Total company sales (pounds and dollars) of all registered product(s) containing the active ingredient(s). If applicable to the active ingredient(s), include foreign sales for those products that are not registered in this country but are applied to sugar (cane or beet), coffee, bananas, cocoa, and other such crops. Present the above information by year for each of the past five years.
- b. Provide an estimate of the sales (pounds and dollars) of the active ingredient(s) for each major use site. Present the above information by year for each of the past five years.
- c. Total direct production cost of product(s) containing the active ingredient(s) by year for the past five years. Include information on raw material cost, direct labor cost, advertising, sales and marketing, and any other significant costs listed separately.
- d. Total indirect production cost (e.g. plant overhead, amortized plant and equipment) charged to product(s) containing the active ingredient(s) by year for the past five years. Exclude all non-recurring costs that were directly related to the active ingredient(s), such as costs of initial registration and any data development.
- e. A list of each data requirement for which you seek a waiver. Indicate the type of waiver sought and the estimated cost to you (listed separately for each data requirement and associated test) of conducting the testing needed to fulfill each of these data requirements.
- f. A list of each data requirement for which you are not seeking any waiver and the estimated cost to you (listed separately for each data requirement and associated test) of conducting the testing needed to fulfill each of these data requirements.
- g. For each of the next ten years, a year-by-year forecast of company sales (pounds and dollars) of the active ingredient(s), direct production costs of product(s) containing the active ingredient(s) (following the parameters in item c above), indirect production costs of product(s)

containing the active ingredient(s) (following the parameters in item d above), and costs of data development pertaining to the active ingredient(s).

- h. A description of the importance and unique benefits of the active ingredient(s) to users. Discuss the use patterns and the effectiveness of the active ingredient(s) relative to registered alternative chemicals and non-chemical control strategies. Focus on benefits unique to the active ingredient(s), providing information that is as quantitative as possible. If you do not have quantitative data upon which to base your estimates, then present the reasoning used to derive your estimates. To assist the Agency in determining the degree of importance of the active ingredient(s) in terms of its benefits, you should provide information on any of the following factors, as applicable to your product(s):
- (1) documentation of the usefulness of the active ingredient(s) in Integrated Pest Management, (b) description of the beneficial impacts on the environment of use of the active ingredient(s), as opposed to its registered alternatives, (c) information on the breakdown of the active ingredient(s) after use and on its persistence in the environment, and (d) description of its usefulness against a pest(s) of public health significance.

Failure to submit sufficient information for the Agency to make a determination regarding a request for a low volume minor use waiver will result in denial of the request for a waiver.

2. Request for Waiver of Data --Option 9 on the Requirements Status and Registrant's Response Form. This option may be used if you believe that a particular data requirement should not apply because the corresponding use is no longer registered or the requirement is inappropriate. You must submit a rationale explaining why you believe the data requirements should not apply. You must also submit the current label(s) of your product(s) and, if a current copy of your Confidential Statement of Formula is not already on file you must submit a current copy.

You will be informed of the Agency's decision in writing. If the Agency determines that the data requirements of this Notice do not apply to your product(s), you will not be required to supply the data pursuant to section 3(c)(2)(B). If EPA determines that the data are required for your product(s), you must choose a method of meeting the requirements of this Notice within the time frame provided by this Notice. Within 30 days of your receipt of the Agency's written decision, you must submit a revised Requirements Status and Registrant's Response Form indicating the option chosen.



# IV. CONSEQUENCES OF FAILURE TO COMPLY WITH THIS NOTICE

# A. NOTICE OF INTENT TO SUSPEND

The Agency may issue a Notice of Intent to Suspend products subject to this Notice due to failure by a registrant to comply with the requirements of this Data Call-In Notice, pursuant to FIFRA section 3(c)(2)(B). Events which may be the basis for issuance of a Notice of Intent to Suspend include, but are not limited to, the following:

- 1. Failure to respond as required by this Notice within 90 days of your receipt of this Notice.
- 2. Failure to submit on the required schedule an acceptable proposed or final protocol when such is required to be submitted to the Agency for review.
- 3. Failure to submit on the required schedule an adequate progress report on a study as required by this Notice.
- 4. Failure to submit on the required schedule acceptable data as required by this Notice.
- 5. Failure to take a required action or submit adequate information pertaining to any option chosen to address the data requirements (e.g., any required action or information pertaining to submission or citation of existing studies or offers, arrangements, or arbitration on the sharing of costs or the formation of Task Forces, failure to comply with the terms of an agreement or arbitration concerning joint data development or failure to comply with any terms of a data waiver).
- 6. Failure to submit supportable certifications as to the conditions of submitted studies, as required by Section III-C of this Notice.
- 7. Withdrawal of an offer to share in the cost of developing required data.
- 8. Failure of the registrant to whom you have tendered an offer to share in the cost of developing data and provided proof of the registrant's receipt of such offer, or failure of a registrant on whom you rely for a generic data exemption either to:
  - a. inform EPA of intent to develop and submit the data required by this Notice on a <u>Data Call-In Response Form</u> and a <u>Requirements Status</u> and <u>Registrant's Response Form</u>; or,
  - b. fulfill the commitment to develop and submit the data as required

by this Notice; or,

- c. otherwise take appropriate steps to meet the requirements stated in this Notice, unless you commit to submit and do submit the required data in the specified time frame.
- 9. Failure to take any required or appropriate steps, not mentioned above, at any time following the issuance of this Notice.

# B. <u>BASIS FOR DETERMINATION THAT SUBMITTED STUDY IS</u> <u>UNACCEPTABLE</u>

The Agency may determine that a study (even if submitted within the required time) is unacceptable and constitutes a basis for issuance of a Notice of Intent to Suspend. The grounds for suspension include, but are not limited to, failure to meet any of the following:

- 1. EPA requirements specified in the Data Call-In Notice or other documents incorporated by reference (including, as applicable, EPA Pesticide Assessment Guidelines, Data Reporting Guidelines, and GeneTox Health Effects Test Guidelines) regarding the design, conduct, and reporting of required studies. Such requirements include, but are not limited to, those relating to test material, test procedures, selection of species, number of animals, sex and distribution of animals, dose and effect levels to be tested or attained, duration of test, and, as applicable, Good Laboratory Practices.
- 2. EPA requirements regarding the submission of protocols, including the incorporation of any changes required by the Agency following review.
- 3. EPA requirements regarding the reporting of data, including the manner of reporting, the completeness of results, and the adequacy of any required supporting (or raw) data, including, but not limited to, requirements referenced or included in this Notice or contained in PR 86-5. All studies must be submitted in the form of a final report; a preliminary report will not be considered to fulfill the submission requirement.

# C. EXISTING STOCKS OF SUSPENDED OR CANCELLED PRODUCTS

EPA has statutory authority to permit continued sale, distribution and use of existing stocks of a pesticide product which has been suspended or cancelled if doing so would be consistent with the purposes of the Federal Insecticide, Fungicide, and Rodenticide Act.



The Agency has determined that such disposition by registrants of existing stocks for a suspended registration when a section 3(c)(2)(B) data request is outstanding would generally not be consistent with the Act's purposes. Accordingly, the Agency anticipates granting registrants permission to sell, distribute, or use existing stocks of suspended product(s) only in exceptional circumstances. If you believe such disposition of existing stocks of your product(s) which may be suspended for failure to comply with this Notice should be permitted, you have the burden of clearly demonstrating to EPA that granting such permission would be consistent with the Act. You must also explain why an "existing stocks" provision is necessary, including a statement of the quantity of existing stocks and your estimate of the time required for their sale, distribution, and use. Unless you meet this burden the Agency will not consider any request pertaining to the continued sale, distribution, or use of your existing stocks after suspension.

If you request a voluntary cancellation of your product(s) as a response to this Notice and your product is in full compliance with all Agency requirements, you will have, under most circumstances, one year from the date your 90 day response to this Notice is due, to sell, distribute, or use existing stocks. Normally, the Agency will allow persons other than the registrant such as independent distributors, retailers and end users to sell, distribute or use such existing stocks until the stocks are exhausted. Any sale, distribution or use of stocks of voluntarily cancelled products containing an active ingredient(s) for which the Agency has particular risk concerns will be determined on case-by-case basis.

Requests for voluntary cancellation received after the 90 day response period required by this Notice will not result in the Agency granting any additional time to sell, distribute, or use existing stocks beyond a year from the date the 90 day response was due unless you demonstrate to the Agency that you are in full compliance with all Agency requirements, including the requirements of this Notice. For example, if you decide to voluntarily cancel your registration six months before a 3 year study is scheduled to be submitted, all progress reports and other information necessary to establish that you have been conducting the study in an acceptable and good faith manner must have been submitted to the Agency, before EPA will consider granting an existing stocks provision.

# SECTION V. <u>REGISTRANTS' OBLIGATION TO REPORT POSSIBLE UNREASONABLE ADVERSE EFFECTS</u>

Registrants are reminded that FIFRA section 6(a)(2) states that if at any time after a pesticide is registered a registrant has additional factual information regarding unreasonable adverse effects on the environment by the pesticide, the registrant shall submit the information to the Agency. Registrants must notify the Agency of any factual information they have, from whatever source, including but not limited to interim or preliminary results of studies, regarding unreasonable adverse effects on man or the environment. This requirement continues as long as the products are registered by the Agency.

# SECTION VI. INOUIRIES AND RESPONSES TO THIS NOTICE

If you have any questions regarding the requirements and procedures established by this Notice, call the contact person listed in Attachment A, the <u>Data Call-In Chemical Status Sheet</u>.

All responses to this Notice (other than voluntary cancellation requests and generic data exemption claims) must include a completed <u>Data Call-In Response Form</u> (Attachment B) and a completed <u>Requirements Status and Registrant's Response Form</u> (Attachment C) and any other documents required by this Notice, and should be submitted to the contact person identified in Attachment A. If the voluntary cancellation or generic data exemption option is chosen, only the <u>Data Call-In Response Form</u> need be submitted.

The Office of Compliance Monitoring (OCM) of the Office of Pesticides and Toxic Substances (OPTS), EPA, will be monitoring the data being generated in response to this Notice.

Sincerely yours,

Peter P. Caulkins Ph.D., Acting Director

Special Review

and Reregistration Division

P. In Carllans

	United
Washington, D.C. 20460	Stales Environmental
).C. 20460	United States Environmental Protection Agency
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1. Company name and Address INSTRUCTIONS: Please type or print in ink. Please read carefully the attached instructions and supply the information requested on this form.
Use additional sheet(s) if necessary "RQUIREMENTS STATUS AND REGISTRANT'S RESPONSE 2. Case # and Name 3. Date and Type of DCI GENERIC OMB No. 2070-0107 2070-0057 Form Approved Approval Expires 03-31-96

10. Certification  I certify that the statements made on this form and all attachments are true, accurate, and complete. I acknowledge that any knowingly false or misleading statement may be punishable by fine, imprisonment or both under applicable law.  Signature and Title of Company's Authorized Representative  12. Name of Company Contact		61-1 Chemical Identity 61-2(a) Begin. mat. & unfg. proc 61-2(b) Discussion of Impurities 62-1 Certification of limits 62-3 Analytical Method	4. Guideline 5. Study Title Requirement Number		
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# Attachment A Chemical Status Sheet

# CEDARWOOD OIL: DATA CALL-IN CHEMICAL STATUS SHEET

### INTRODUCTION

You have been sent this Generic Data Call-In Notice because you have a product containing cedarwood oil.

This Generic Data Call-In Chemical Status Sheet, contains an overview of data required by this notice, and a point of contact for inquiries pertaining to the reregistration of cedarwood oil. This attachment is to be used in conjunction with (1) the Generic Data Call-In Notice, (2) the Generic Data Call-In Response Form (Attachment B), (3) the Requirements Status and Registrant's Form (Attachment C), (4) a list of registrants receiving this DCI (Attachment D), (5) the EPA Acceptance Criteria (Attachment E), and (6) the Cost Share and Data Compensation Forms in replying to this cedarwood oil Generic Data Call-In (Attachment F). Instructions and guidance accompany each form.

# DATA REQUIRED BY THIS NOTICE

The additional data requirements needed to complete the generic database for cedarwood oil are contained in the Requirements Status and Registrant's Response, Attachment C. The Agency has concluded that additional product chemistry data on cedarwood oil needed. These data are needed to fully complete the reregistration of all eligible cedarwood oil products.

# INQUIRIES AND RESPONSES TO THIS NOTICE

If you have any questions regarding the generic data requirements and procedures established by this Notice, please contact Virginia Dietrich at (703) 308-8157. All responses to this Notice for the generic data requirements should be submitted to:

> Virginia Dietrich, Chemical Review Manager Accelerated Reregistration Branch Special Review and Registration Division (7508W) Office of Pesticide Programs U.S. Environmental Protection Agency Washington, D.C. 20460

-- RE: -- Cedarwood Oil



# Attachment B Generic DCI Response Forms (Form A) plus Instructions

### SPECIFIC INSTRUCTIONS FOR THE DATA CALL-IN RESPONSE FORM

This Form is designed to be used to respond to call-ins for generic and product specific data for the purpose of reregistering pesticides under the Federal Insecticide Fungicide and Rodenticide Act. Fill out this form each time you are responding to a data call-in for which EPA has sent you the form entitled "Requirements Status and Registrant's Response."

Items 1-4 will have been preprinted on the form Items 5 through 7 must be completed by the registrant as appropriate Items 8 through 11 must be completed by the registrant before submitting a response to the Agency.

Public reporting burden for this collection of information is estimated to average 15 minutes per response, including time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding the burden estimate or any other aspect of this collection of information, including suggesting for reducing this burden, to Chief, Information Policy Branch, PM-223, U S Environmental Protection Agency, 401 M St, S W, Washington, D C 20460; and to the Office of Management and Budget, Paperwork Reduction Project 2070-0107, Washington, D C 20503

#### INSTRUCTIONS

- Item 1 This item identifies your company name, number and address.
- Item 2 This item identifies the ease number, ease name, EPA chemical number and chemical name.
- Item 3 This item identifies the date and type of data call-in.
- Item 4 This item identifies the EPA product registrations relevant to the data call-in. Please note that you are also responsible for informing the Agency of your response regarding any product that you believe may be covered by this data call-in but that is not listed by the Agency in Item 4. You must bring any such apparent omission to the Agency's attention within the period required for submission of this response form.
- Item 5 Cheek this item for each product registration you wish to cancel voluntarily. If a registration number is listed for a product for which you previously requested voluntary cancellation, indicate in Item 5 the date of that request. You do not need to complete any item on the Requirements Status and Registrant's Response Form for any product that is voluntarily cancelled.
- Item 6a Check this item if this data call-in is for generic data as indicated in Item 3 and if you are eligible for a Generic Data Exemption for the chemical listed in Item

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2 and used in the subject product. By electing this exemption, you agree to the terms and conditions of a Generic Data Exemption as explained in the Data Call-In Notice.

If you are eligible for or claim a Generic Data Exemption, enter the EPA registration Number of each registered source of that active ingredient that you use in your product.

Typically, if you purchase an EPA-registered product from one or more other producers (who, with respect to the incorporated product, are in compliance with this and-any other outstanding Data Call-In Notice), and incorporate that product into all your products, you may complete this item for all products listed on this form If, however, you produce the active ingredient yourself, or use any unregistered product (regardless of the fact that some of your sources are registered), you may not claim a Generic Data Exemption and you may not select this item.

- Item 6b Check this Item if the data call-in is a generic data call-in as indicated in Item 3 and if you are agreeing to satisfy the generic data requirements of this data call-in. Attach the <u>Requirements Status and Registrant's Response Form</u> that Indicates how you will satisfy those requirements.
- Item 7a Check this item only if this call-in is a data call-in as indicated in Item 3 for a manufacturing use product (MUP), and if your product is a manufacturing use product for which you agree to supply product-specific data. Attach the Requirements Status and Registrants' Response Form that indicates how you will satisfy those requirements.
- Check this item only if this call-in is a data call-in for an end use product (EUP) as indicated in Item 3 and if your product is an end use product for which you agree to supply product-specific data. Attach the Requirements Status and Registrant's Response Form that indicates how you will satisfy those requirements.
- Item 8 This certification statement must be signed by an authorized representative of your company and the person signing must include his/her title. Additional pages used in your response must be initialled and dated in the space provided for the certification.
- Item 9 Enter the date of signature.
- Item 10 Enter the name of the person EPA should contact with questions regarding your response.

Item 11 Enter the phone number of your company contact.

# Attachment C

Requirements Status and Registrants' Response Forms (Form B) plus Instructions

# SPECIFIC INSTRUCTIONS FOR COMPLETING THE REQUIREMENTS STATUS AND REGISTRANTS RESPONSE FORM

#### Generic Data

This form is designed to be used for registrants to respond to call-in- for generic and product-specific data as part of EPA's reregistration program under the Federal Insecticide Fungicide and Rodenticide Act. Although the <u>form</u> is the same for both product specific and generic data, <u>instructions</u> for completing the forms differ slightly. Specifically, options for satisfying product specific data requirements do not include (1) deletion of uses or (2) request for a low volume/minor use waiver. These instructions are for completion of generic data requirements.

EPA has developed this form individually for each data call-in addressed to each registrant, and has preprinted this form with a number of items. **DO NOT** use this form for any other active ingredient.

Items 1 through 8 (inclusive) will have been preprinted on the form. You must complete all other items on this form by typing or printing legibly.

Public reporting burden for this collection of information is estimated to average 30 minutes per response, including time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding the burden estimate or any other aspect of this collection of information, including suggesting for reducing this burden, to Chief, Information Policy Branch, PM-223, U.S. Environmental Protection Agency, 401 M St., S.W., Washington, D.C. 20460; and to the Office of Management and Budget, Paperwork Reduction Project 2070-0107, Washington, D.C. 20503.

### **INSTRUCTIONS**

- Item 1. This item identifies your company name, number, and address.
- Item 2. This item identifies the case number, case name, EPA chemical number and chemical name.
- Item 3. This item identifies the date and type of data call-in.
- Item 4. This item identifies the guideline reference numbers of studies required to support the product(s) being reregistered. These guidelines, in addition to requirements specified in the Data Call-In Notice, govern the conduct of the required studies.
- Item 5. This item identifies the study title associated with the guideline reference



number and whether protocols and 1, 2, or 3-year progress reports are required to be submitted in connection with the study. As noted in Section III of the Data Call-In Notice, 90-day progress reports are required for all studies.

If an asterisk appears in Item 5, EPA has attached information relevant to this guideline reference number to the Requirements Status and Registrant's Response Form.

Item 6. This item identifies the code associated with the use pattern of the pesticide. A brief description of each code follows:

A.	Terrestrial food
B.	Terrestrial feed
C.	Terrestrial non-food
D.	Aquatic food
E.	Aquatic non-food outdoor
F.	Aquatic non-food industrial
G.	Aquatic non-food residential
H.	Greenhouse food
I.	Greenhouse non-food crop
J.	Forestry
K.	Residential
L.	Indoor food
M.	Indoor non-food
N.	Indoor medical
O	Indoor residential

Item 7. This item identifies the code assigned to the substance that must be used for testing. A brief description of each code follows.

EP	End-Use Product
MP	Manufacturing-Use Product
MP/TGAI	Manufacturing-Use Product and Technical Grade Active
	Ingredient
PAI	Pure Active Ingredient
PAI/M	Pure Active Ingredient and Metabolites
PAI/PAIRA	Pure Active Ingredient or Pure Active Ingredient
	Radiolabelled
PAIRA	Pure Active Ingredient Radiolabelled
PAIRA/M	Pure Active Ingredient Radiolabelled and Metabolites
PAIRA/PM	Pure Active Ingredient Radiolabelled and Plant
	Metabolites
TEP	Typical End-Use Product
TEP *	Typical End-Use Product, Percent Active Ingredient

Specified

TEP/MET Typical End-Use Product and Metabolites

TEP/PAI/M Typical End-Use Product or Pure Active Ingredient and

Metabolites

TGAI/PAIRA Technical Grade Active Ingredient or Pure Active

Ingredient Radiolabelled

TGAI Technical Grade Active Ingredient

TGAI/TEP Technical Grade Active Ingredient or Typical

End-Use Product

TGAI/PAI Technical Grade Active Ingredient or Pure Active

Ingredient

MET Metabolites IMP Impurities

DEGR Degradates

\*See: guideline comment

Item 8. This item identifies the time frame allowed for submission of the study or protocol identified in item 2. The time frame runs from the date of your receipt of the Data Call-In Notice.

- Item 9. Enter the appropriate Response Code or Codes to show how you intend to comply with each data requirement. Brief descriptions of each code follow.

  The Data Call-In Notice contains a fuller description of each of these options.
  - 1. (Developing Data) I will conduct a new study and submit it within the time frames specified in item 8 above. By indicating that I have chosen this option, I certify that I will comply with all the requirements pertaining to the conditions for submittal of this study as outlined in the Data Call-In Notice and that I will provide the protocol and progress reports required in item 5 above.
  - 2. (Agreement to Cost Share) I have entered into an agreement with one or more registrants to develop data jointly. By indicating that I have chosen this option, I certify that I will comply with all the requirements pertaining to sharing in the cost of developing data as outlined in the Data Call-ln Notice.
  - 3. (Offer to Cost Share) I have made an offer to enter into an agreement with one or more registrants to develop data jointly. I am submitting a copy of the form "Certification of Offer to Cost Share in the Development of Data" that describes this offer/agreement. By indicating that I have chosen this option, I certify that I will comply with all the requirements pertaining to making an offer to share in the

cost of developing data as outlined in the Data Call-In Notice.

- 4. (Submitting Existing Data) I am submitting an existing study that has never before been submitted to EPA. By indicating that I have chosen this option, I certify that this study meets all the requirements pertaining to the conditions for submittal of existing data outlined in the Data Call-In Notice and I have attached the needed supporting information along with this response.
- 5. (Upgrading a Study) I am submitting or citing data to upgrade a study that EPA has classified as partially acceptable and potentially upgradeable. By indicating that I have chosen this option, I certify that I have met all the requirements pertaining to the conditions for submitting or citing existing data to upgrade a study described in the Data Call-In Notice. I am indicating on attached correspondence the Master Record Identification Number (MRID) that EPA has assigned to the data that I am citing as well as the MRID of the study I am attempting to upgrade.
- 6. (Citing a Study) I am citing an existing study that has been previously classified by EPA as acceptable, core, core minimum, or a study that has not yet been reviewed by the Agency. I am providing the Agency's classification of the study.
- 7. (Deleting Uses) I am attaching an application for amendment to my registration deleting the uses for which the data are required.
- 8. (Low Volume/Minor Use Waiver Request) I have read the statements concerning low volume-minor use data waivers in the Data Call-In Notice and I request a low-volume minor use waiver of the data requirement. I am attaching a detailed justification to support this waiver request including, among other things, all information required to support the request. I understand that, unless modified by the Agency in writing, the data requirement as stated in the Notice governs.
- 9. (Request for Waiver of Data) I have read the statements concerning data waivers other than low volume minor-use data waivers in the Data Call-In Notice and I request a waiver of the data requirement. I am attaching an identification of the basis for this waiver and a detailed justification to support this waiver request. The justification includes, among other things, all information required to support the request. I understand that, unless modified by the Agency in writing, the data requirement as stated in the Notice governs.

- Item 10. This item must be signed by an authorized representative of your company. The person signing must include his/her title, and must initial and date all other pages of this form.
- Item 11. Enter the date of signature.
- Item 12. Enter the name of the person EPA should contact with questions regarding your response.
- Item 13. Enter the phone number of your company contact.

# Attachment D List of Registrants Sent This DCI

## APPENDIX G

Product Specific Data Call-In

#### DATA CALL-IN NOTICE

### CERTIFIED MAIL

#### Dear Sir or Madam:

This Notice requires you and other registrants of pesticide products containing the active ingredient identified in Attachment A of this Notice, the <u>Data Call-In Chemical Status Sheet</u>, to submit certain product specific data as noted herein to the U.S. Environmental Protection Agency (EPA, the Agency). These data are necessary to maintain the continued registration of your product(s) containing this active ingredient. Within 90 days after you receive this Notice you must respond as set forth in Section III below. Your response must state:

- 1. How you will comply with the requirements set forth in this Notice and its Attachments A through G; or
- 2. Why you believe you are exempt from the requirements listed in this Notice and in Attachment C, Requirements Status and Registrant's Response Form, (see section III-B); or
- 3. Why you believe EPA should not require your submission of product specific data in the manner specified by this Notice (see section III-D).

If you do not respond to this Notice, or if you do not satisfy EPA that you will comply with its requirements or should be exempt or excused from doing so, then the registration of your product(s) subject to this Notice will be subject to suspension. We have provided a list of all of your products subject to this Notice in Attachment B, <u>Data Call-In Response Form</u>, as well as a list of all registrants who were sent this Notice (Attachment F).

The authority for this Notice is section 3(c)(2)(B) of the Federal Insecticide, Fungicide and Rodenticide Act as amended (FIFRA), 7 U.S.C. section 136a(c)(2)(B). Collection of this information is authorized under the Paperwork Reduction Act by OMB Approval No. 2070-0107 (expiration date 12-31-92).



This Notice is divided into six sections and seven Attachments. The Notice itself contains information and instructions applicable to all Data Call-In Notices. The Attachments contain specific chemical information and instructions. The six sections of the Notice are:

Section I - Why You Are Receiving This Notice

Section II - Data Required By This Notice

Section III- Compliance With Requirements Of This Notice

Section IV - Consequences Of Failure To Comply With This Notice

Section V - Registrants' Obligation To Report Possible Unreasonable

Adverse Effects

Section VI - Inquiries And Responses To This Notice

#### The Attachments to this Notice are:

- A Data Call-In Chemical Status Sheet
- B Product-Specific Data Call-In Response Form
- C Requirements Status and Registrant's Response Form
- D EPA Grouping of End-Use Products for Meeting Acute Toxicology Data Requirements for Reregistration
- E EPA Acceptance Criteria
- F List of Registrants Receiving This Notice
- G Cost Share and Data Compensation Forms, and Product Specific Data Report
  Form

#### SECTION I. WHY YOU ARE RECEIVING THIS NOTICE

The Agency has reviewed existing data for this active ingredient and reevaluated the data needed to support continued registration of the subject active ingredient. The Agency has concluded that the only additional data necessary are product specific data. No additional generic data requirements are being imposed. You have been sent this Notice because you have product(s) containing the subject active ingredient.

### SECTION II. DATA REQUIRED BY THIS NOTICE

#### II-A. DATA REQUIRED

The product specific data required by this Notice are specified in Attachment C, Requirements Status and Registrant's Response Form. Depending on the results of the studies required in this Notice, additional testing may be required.



### II-B. SCHEDULE FOR SUBMISSION OF DATA

You are required to submit the data or otherwise satisfy the data requirements specified in Attachment C, Requirements Status and Registrant's Response Form, within the time frames provided.

#### II-C. TESTING PROTOCOL

All studies required under this Notice must be conducted in accordance with test standards outlined in the Pesticide Assessment Guidelines for those studies for which guidelines have been established.

These EPA Guidelines are available from the National Technical Information Service (NTIS), Attn: Order Desk, 5285 Port Royal Road, Springfield, Va 22161 (tel: 703-487-4650).

Protocols approved by the Organization for Economic Cooperation and Development (OECD) are also acceptable if the OECD-recommended test standards conform to those specified in the Pesticide Data Requirements regulation (40 CFR § 158.70). When using the OECD protocols, they should be modified as appropriate so that the data generated by the study will satisfy the requirements of 40 CFR § 158. Normally, the Agency will not extend deadlines for complying with data requirements when the studies were not conducted in accordance with acceptable standards. The OECD protocols are available from OECD, 1750 Pennsylvania Avenue N.W., Washington, D.C. 20006.

All new studies and proposed protocols submitted in response to this Data Call-In Notice must be in accordance with Good Laboratory Practices [40 CFR Part 160.3(a)(6)].

# II-D. <u>REGISTRANTS RECEIVING PREVIOUS SECTION 3(c)(2)(B) NOTICES</u> ISSUED BY THE AGENCY

Unless otherwise noted herein, this Data Call-In does not in any way supersede or change the requirements of any previous Data Call-In(s), or any other agreements entered into with the Agency pertaining to such prior Notice. Registrants must comply with the requirements of all Notices to avoid issuance of a Notice of Intent to Suspend their affected products.

#### SECTION III. COMPLIANCE WITH REQUIREMENTS OF THIS NOTICE

#### III-A. SCHEDULE FOR RESPONDING TO THE AGENCY

The appropriate responses initially required by this Notice for product specific data must be submitted to the Agency within 90 days after your receipt of this Notice. Failure to adequately respond to this Notice within 90 days of your receipt will be a basis for issuing a Notice of Intent to Suspend (NOIS) affecting your products. This and other bases for issuance



of NOIS due to failure to comply with this Notice are presented in Section IV-A and IV-B.

#### III-B. OPTIONS FOR RESPONDING TO THE AGENCY

The options for responding to this Notice for product specific data are: (a) voluntary cancellation, (b) agree to satisfy the product specific data requirements imposed by this notice or (c) request a data waiver(s).

A discussion of how to respond if you chose the Voluntary Cancellation option is presented below. A discussion of the various options available for satisfying the product specific data requirements of this Notice is contained in Section III-C. A discussion of options relating to requests for data waivers is contained in Section III-D.

There are two forms that accompany this Notice of which, depending upon your response, one or both must be used in your response to the Agency. These forms are the <u>Data-Call-In Response Form</u>, and the <u>Requirements Status and Registrant's Response Form</u>, Attachment B and Attachment C. The <u>Data Call-In Response Form</u> must be submitted as part of every response to this Notice. In addition, one copy of the <u>Requirements Status and Registrant's Response Form</u> must be submitted for each product listed on the <u>Data Call-In Response Form</u> unless the voluntary cancellation option is selected or unless the product is identical to another (refer to the instructions for completing the <u>Data Call-In Response Form</u> in Attachment B). Please note that the company's authorized representative is required to sign the first page of the <u>Data Call-In Response Form</u> and <u>Requirements Status and Registrant's Response Form</u> (if this form is required) and initial any subsequent pages. The forms contain separate detailed instructions on the response options. Do not alter the printed material. If you have questions or need assistance in preparing your response, call or write the contact person(s) identified in Attachment A.

1. Voluntary Cancellation - You may avoid the requirements of this Notice by requesting voluntary cancellation of your product(s) containing the active ingredient that is the subject of this Notice. If you wish to voluntarily cancel your product, you must submit a completed <u>Data Call-In Response Form</u>, indicating your election of this option. Voluntary cancellation is item number 5 on the <u>Data Call-In Response Form</u>. If you choose this option, this is the only form that you are required to complete.

If you chose to voluntarily cancel your product, further sale and distribution of your product after the effective date of cancellation must be in accordance with the Existing Stocks provisions of this Notice which are contained in Section IV-C.

2. Satisfying the Product Specific Data Requirements of this Notice There are various options available to satisfy the product specific data requirements of this Notice. These options are discussed in Section III-C of this Notice and comprise options 1 through 6 on the



Requirements Status and Registrant's Response Form and item numbers 7a and 7b on the <u>Data Call-In Response Form</u>. Deletion of a use(s) and the low volume/minor use option are not valid options for fulfilling product specific data requirements.

3. Request for Product Specific Data Waivers. Waivers for product specific data are discussed in Section III-D of this Notice and are covered by option 7 on the Requirements Status and Registrant's Response Form. If you choose one of these options, you must submit both forms as well as any other information/data pertaining to the option chosen to address the data requirement.

#### III-C SATISFYING THE DATA REQUIREMENTS OF THIS NOTICE

If you acknowledge on the <u>Data Call-In Response Form</u> that you agree to satisfy the product specific data requirements (i.e. you select item number 7a or 7b), then you must select one of the six options on the <u>Requirements Status and Registrant's Response Form</u> related to data production for each data requirement. Your option selection should be entered under item number 9, "Registrant Response." The six options related to data production are the first six options discussed under item 9 in the instructions for completing the <u>Requirements Status and Registrant's Response Form</u>. These six options are listed immediately below with information in parentheses to guide registrants to additional instructions provided in this Section. The options are:

- (1) I will generate and submit data within the specified time frame (Developing Data)
- (2) I have entered into an agreement with one or more registrants to develop data jointly (Cost Sharing)
- (3) I have made offers to cost-share (Offers to Cost Share)
- (4) I am submitting an existing study that has not been submitted previously to the Agency by anyone (Submitting an Existing Study)
- (5) I am submitting or citing data to upgrade a study classified by EPA as partially acceptable and upgradeable (Upgrading a Study)
- (6) I am citing an existing study that EPA has classified as acceptable or an existing study that has been submitted but not reviewed by the Agency (Citing an Existing Study)

Option 1, Developing Data -- If you choose to develop the required data it must be in conformance with Agency deadlines and with other Agency requirements as referenced herein and in the attachments. All data generated and submitted must comply with the Good Laboratory Practice (GLP) rule (40 CFR Part 160), be conducted according to the Pesticide Assessment Guidelines (PAG), and be in conformance with the requirements of PR Notice 86-5.

The time frames in the <u>Requirements Status and Registrant's Response Form</u> are the time frames that the Agency is allowing for the submission of completed study reports. The noted deadlines run from the date of the receipt of this Notice by the registrant. If the data are not



submitted by the deadline, each registrant is subject to receipt of a Notice of Intent to Suspend the affected registration(s).

If you cannot submit the data/reports to the Agency in the time required by this Notice and intend to seek additional time to meet the requirements(s), you must submit a request to the Agency which includes: (1) a detailed description of the expected difficulty and (2) a proposed schedule including alternative dates for meeting such requirements on a step-by-step basis. You must explain any technical or laboratory difficulties and provide documentation from the laboratory performing the testing. While EPA is considering your request, the original deadline remains. The Agency will respond to your request in writing. If EPA does not grant your request, the original deadline remains. Normally, extensions can be requested only in cases of extraordinary testing problems beyond the expectation or control of the registrant. Extensions will not be given in submitting the 90-day responses. Extensions will not be considered if the request for extension is not made in a timely fashion; in no event shall an extension request be considered if it is submitted at or after the lapse of the subject deadline.

Option 2. Agreement to Share in Cost to Develop Data -- Registrants may only choose this option for acute toxicity data and certain efficacy data and only if EPA has indicated in the attached data tables that your product and at least one other product are similar for purposes of depending on the same data. If this is the case, data may be generated for just one of the products in the group. The registration number of the product for which data will be submitted must be noted in the agreement to cost share by the registrant selecting this option. If you choose to enter into an agreement to share in the cost of producing the required data but will not be submitting the data yourself, you must provide the name of the registrant who will be submitting the data. You must also provide EPA with documentary evidence that an agreement has been formed. Such evidence may be your letter offering to join in an agreement and the other registrant's acceptance of your offer, or a written statement by the parties that an agreement exists. The agreement to produce the data need not specify all of the terms of the final arrangement between the parties or the mechanism to resolve the terms. Section 3(c)(2)(B) provides that if the parties cannot resolve the terms of the agreement they may resolve their differences through binding arbitration.

Option 3. Offer to Share in the Cost of Data Development -- This option only applies to acute toxicity and certain efficacy data as described in option 2 above. If you have made an offer to pay in an attempt to enter into an agreement or amend an existing agreement to meet the requirements of this Notice and have been unsuccessful, you may request EPA (by selecting this option) to exercise its discretion not to suspend your registration(s), although you do not comply with the data submission requirements of this Notice. EPA has determined that as a general policy, absent other relevant considerations, it will not suspend the registration of a product of a registrant who has in good faith sought and continues to seek to enter into a joint data development/cost sharing program, but the other registrant(s) developing the data has refused to accept your offer. To qualify for this option, you must submit documentation to the Agency proving that you have made an offer to another registrant (who has an obligation to submit data) to share in the burden of developing that data. You must also submit to the Agency a completed EPA Form 8570-32, Certification of Offer to Cost Share in the Development of Data, Attachment G. In addition, you must demonstrate that the other registrant to whom the offer was made has not accepted your offer to enter into a cost sharing

agreement by including a copy of your offer and proof of the other registrant's receipt of that offer (such as a certified mail receipt). Your offer must, in addition to anything else, offer to share in the burden of producing the data upon terms to be agreed or failing agreement to be bound by binding arbitration as provided by FIFRA section 3(c)(2)(B)(iii) and must not qualify this offer. The other registrant must also inform EPA of its election of an option to develop and submit the data required by this Notice by submitting a <u>Data Call-In Response Form</u> and a <u>Requirements Status and Registrant's Response Form</u> committing to develop and submit the data required by this Notice.

In order for you to avoid suspension under this option, you may not withdraw your offer to share in the burdens of developing the data. In addition, the other registrant must fulfill its commitment to develop and submit the data as required by this Notice. If the other registrant fails to develop the data or for some other reason is subject to suspension, your registration as well as that of the other registrant will normally be subject to initiation of suspension proceedings, unless you commit to submit, and do submit the required data in the specified time frame. In such cases, the Agency generally will not grant a time extension for submitting the data.

Option 4, Submitting an Existing Study -- If you choose to submit an existing study in response to this Notice, you must determine that the study satisfies the requirements imposed by this Notice. You may only submit a study that has not been previously submitted to the Agency or previously cited by anyone. Existing studies are studies which predate issuance of this Notice. Do not use this option if you are submitting data to upgrade a study. (See Option 5).

You should be aware that if the Agency determines that the study is not acceptable, the Agency will require you to comply with this Notice, normally without an extension of the required date of submission. The Agency may determine at any time that a study is not valid and needs to be repeated.

To meet the requirements of the DCI Notice for submitting an existing study, all of the following three criteria must be clearly met:

a. You must certify at the time that the existing study is submitted that the raw data and specimens from the study are available for audit and review and you must identify where they are available. This must be done in accordance with the requirements of the Good Laboratory Practice (GLP) regulation, 40 CFR Part 160. As stated in 40 CFR 160.3(j) " 'raw data' means any laboratory worksheets, records, memoranda, notes, or exact copies thereof, that are the result of original observations and activities of a study and are necessary for the reconstruction and evaluation of the report of that study. In the event that exact transcripts of raw data have been prepared (e.g., tapes which have been transcribed verbatim, dated, and verified accurate by signature), the exact copy or exact transcript may be substituted for the original source as raw data. 'Raw data' may include photographs, microfilm or microfiche copies, computer printouts, magnetic



media, including dictated observations, and recorded data from automated instruments." The term "specimens", according to 40 CFR 160.3(k), means "any material derived from a test system for examination or analysis."

- b. Health and safety studies completed after May 1984 must also contain all GLP-required quality assurance and quality control information, pursuant to the requirements of 40 CFR Part 160. Registrants must also certify at the time of submitting the existing study that such GLP information is available for post-May 1984 studies by including an appropriate statement on or attached to the study signed by an authorized official or representative of the registrant.
- c. You must certify that each study fulfills the acceptance criteria for the Guideline relevant to the study provided in the FIFRA Accelerated Reregistration Phase 3 Technical Guidance and that the study has been conducted according to the Pesticide Assessment Guidelines (PAG) or meets the purpose of the PAG (both available from NTIS). A study not conducted according to the PAG may be submitted to the Agency for consideration if the registrant believes that the study clearly meets the purpose of the PAG. The registrant is referred to 40 CFR 158.70 which states the Agency's policy regarding acceptable protocols. If you wish to submit the study, you must, in addition to certifying that the purposes of the PAG are met by the study, clearly articulate the rationale why you believe the study meets the purpose of the PAG, including copies of any supporting information or data. It has been the Agency's experience that studies completed prior to January 1970 rarely satisfied the purpose of the PAG and that necessary raw data are usually not available for such studies.

If you submit an existing study, you must certify that the study meets all requirements of the criteria outlined above.

If you know of a study pertaining to any requirement in this Notice which does not meet the criteria outlined above but does contain factual information regarding unreasonable adverse effects, you must notify the Agency of such a study. If such study is in the Agency's files, you need only cite it along with the notification. If not in the Agency's files, you must submit a summary and copies as required by PR Notice 86-5.

Option 5, Upgrading a Study -- If a study has been classified as partially acceptable and upgradeable, you may submit data to upgrade that study. The Agency will review the data submitted and determine if the requirement is satisfied. If the Agency decides the requirement is not satisfied, you may still be required to submit new data normally without any time extension. Deficient, but upgradeable studies will normally be classified as supplemental. However, it is important to note that not all studies classified as supplemental are upgradeable. If you have questions regarding the classification of a study or whether a study may be upgraded, call or write the contact person listed in Attachment A. If you submit data to



upgrade an existing study you must satisfy or supply information to correct <u>all</u> deficiencies in the study identified by EPA. You must provide a clearly articulated rationale of how the deficiencies have been remedied or corrected and why the study should be rated as acceptable to EPA. Your submission must also specify the MRID number(s) of the study which you are attempting to upgrade and must be in conformance with PR Notice 86-5.

Do not submit additional data for the purpose of upgrading a study classified as unacceptable and determined by the Agency as not capable of being upgraded.

This option should also be used to cite data that has been previously submitted to upgrade a study, but has not yet been reviewed by the Agency. You must provide the MRID number of the data submission as well as the MRID number of the study being upgraded.

The criteria for submitting an existing study, as specified in Option 4 above, apply to all data submissions intended to upgrade studies. Additionally your submission of data intended to upgrade studies must be accompanied by a certification that you comply with each of those criteria as well as a certification regarding protocol compliance with Agency requirements.

Option 6, Citing Existing Studies -- If you choose to cite a study that has been previously submitted to EPA, that study must have been previously classified by EPA as acceptable or it must be a study which has not yet been reviewed by the Agency. Acceptable toxicology studies generally will have been classified as "core-guideline" or "core minimum." For all other disciplines the classification would be "acceptable." With respect to any studies for which you wish to select this option you must provide the MRID number of the study you are citing and, if the study has been reviewed by the Agency, you must provide the Agency's classification of the study.

If you are citing a study of which you are not the original data submitter, you must submit a completed copy of EPA Form 8570-31, Certification with Respect to Data Compensation Requirements.

Registrants who select one of the above 6 options must meet all of the requirements described in the instructions for completing the <u>Data Call-In Response</u> Form and the <u>Requirements Status and Registrant's Response</u> Form, as appropriate.

#### III-D REQUESTS FOR DATA WAIVERS

If you request a waiver for product specific data because you believe it is inappropriate, you must attach a complete justification for the request, including technical reasons, data and references to relevant EPA regulations, guidelines or policies. (Note: any supplemental data must be submitted in the format required by PR Notice 86-5). This will be the only opportunity to state the reasons or provide information in support of your request. If the Agency approves your waiver request, you will not be required to supply the data pursuant to section 3(c)(2)(B) of FIFRA. If the Agency denies your waiver request, you must choose an option for meeting the data requirements of this Notice within 30 days of the receipt of the Agency's decision. You must indicate and submit the option chosen on the Requirements Status and Registrant's Response Form. Product specific data requirements for product chemistry, acute toxicity and efficacy (where appropriate) are required for all products and the Agency would grant a waiver only under extraordinary circumstances. You should also be aware that submitting a waiver request will not automatically extend the due date for the study in question.

Waiver requests submitted without adequate supporting rationale will be denied and the original due date will remain in force.

#### IV. CONSEQUENCES OF FAILURE TO COMPLY WITH THIS NOTICE

#### IV-A NOTICE OF INTENT TO SUSPEND

The Agency may issue a Notice of Intent to Suspend products subject to this Notice due to failure by a registrant to comply with the requirements of this Data Call-In Notice, pursuant to FIFRA section 3(c)(2)(B). Events which may be the basis for issuance of a Notice of Intent to Suspend include, but are not limited to, the following:

- 1. Failure to respond as required by this Notice within 90 days of your receipt of this Notice.
- 2. Failure to submit on the required schedule an acceptable proposed or final protocol when such is required to be submitted to the Agency for review.
- 3. Failure to submit on the required schedule an adequate progress report on a study as required by this Notice.
- 4. Failure to submit on the required schedule acceptable data as required by this Notice.
- 5. Failure to take a required action or submit adequate information pertaining to any option chosen to address the data requirements (e.g., any required action or information pertaining to submission or citation of existing studies or offers, arrangements, or arbitration on the sharing of costs or the formation of Task Forces, failure to comply with the terms of an agreement or arbitration concerning joint data development or failure to comply with any terms of a data waiver).
- 6. Failure to submit supportable certifications as to the conditions of submitted studies, as required by Section III-C of this Notice.
- 7. Withdrawal of an offer to share in the cost of developing required data.
- 8. Failure of the registrant to whom you have tendered an offer to share in the cost of developing data and provided proof of the registrant's receipt of such offer or failure of a registrant on whom you rely for a generic data exemption either to:
  - a. inform EPA of intent to develop and submit the data required by this Notice on a <u>Data Call-In Response Form</u> and a <u>Requirements Status and Registrant's Response Form</u>;
  - b. fulfill the commitment to develop and submit the data as required by this Notice; or



- c. otherwise take appropriate steps to meet the requirements stated in this Notice, unless you commit to submit and do submit the required data in the specified time frame.
- 9. Failure to take any required or appropriate steps, not mentioned above, at any time following the issuance of this Notice.

# IV-B. BASIS FOR DETERMINATION THAT SUBMITTED STUDY IS UNACCEPTABLE

The Agency may determine that a study (even if submitted within the required time) is unacceptable and constitutes a basis for issuance of a Notice of Intent to Suspend. The grounds for suspension include, but are not limited to, failure to meet any of the following:

- 1. EPA requirements specified in the Data Call-In Notice or other documents incorporated by reference (including, as applicable, EPA Pesticide Assessment Guidelines, Data Reporting Guidelines, and GeneTox Health Effects Test Guidelines) regarding the design, conduct, and reporting of required studies. Such requirements include, but are not limited to, those relating to test material, test procedures, selection of species, number of animals, sex and distribution of animals, dose and effect levels to be tested or attained, duration of test, and, as applicable, Good Laboratory Practices.
- 2. EPA requirements regarding the submission of protocols, including the incorporation of any changes required by the Agency following review.
- 3. EPA requirements regarding the reporting of data, including the manner of reporting, the completeness of results, and the adequacy of any required supporting (or raw) data, including, but not limited to, requirements referenced or included in this Notice or contained in PR 86-5. All studies must be submitted in the form of a final report; a preliminary report will not be considered to fulfill the submission requirement.

#### IV-C EXISTING STOCKS OF SUSPENDED OR CANCELLED PRODUCTS

EPA has statutory authority to permit continued sale, distribution and use of existing stocks of a pesticide product which has been suspended or cancelled if doing so would be consistent with the purposes of the Act.

The Agency has determined that such disposition by registrants of existing stocks for a suspended registration when a section 3(c)(2)(B) data request is outstanding would generally not be consistent with the Act's purposes. Accordingly, the Agency anticipates granting registrants permission to sell, distribute, or use existing stocks of suspended product(s) only in exceptional circumstances. If you believe such disposition of existing stocks of your product(s) which may be suspended for failure to comply with this Notice should be permitted, you have the burden of clearly demonstrating to EPA that granting such permission would be consistent with the Act.



You must also explain why an "existing stocks" provision is necessary, including a statement of the quantity of existing stocks and your estimate of the time required for their sale, distribution, and use. Unless you meet this burden the Agency will not consider any request pertaining to the continued sale, distribution, or use of your existing stocks after suspension.

If you request a voluntary cancellation of your product(s) as a response to this Notice and your product is in full compliance with all Agency requirements, you will have, under most circumstances, one year from the date your 90 day response to this Notice is due, to sell, distribute, or use existing stocks. Normally, the Agency will allow persons other than the registrant such as independent distributors, retailers and end users to sell, distribute or use such existing stocks until the stocks are exhausted. Any sale, distribution or use of stocks of voluntarily cancelled products containing an active ingredient for which the Agency has particular risk concerns will be determined on case-by-case basis.

Requests for voluntary cancellation received after the 90 day response period required by this Notice will not result in the Agency granting any additional time to sell, distribute, or use existing stocks beyond a year from the date the 90 day response was due unless you demonstrate to the Agency that you are in full compliance with all Agency requirements, including the requirements of this Notice. For example, if you decide to voluntarily cancel your registration six months before a 3 year study is scheduled to be submitted, all progress reports and other information necessary to establish that you have been conducting the study in an acceptable and good faith manner must have been submitted to the Agency, before EPA will consider granting an existing stocks provision.

# SECTION V. <u>REGISTRANTS' OBLIGATION TO REPORT POSSIBLE</u> UNREASONABLE ADVERSE EFFECTS

Registrants are reminded that FIFRA section 6(a)(2) states that if at any time after a pesticide is registered a registrant has additional factual information regarding unreasonable adverse effects on the environment by the pesticide, the registrant shall submit the information to the Agency. Registrants must notify the Agency of any factual information they have, from whatever source, including but not limited to interim or preliminary results of studies, regarding unreasonable adverse effects on man or the environment. This requirement continues as long as the products are registered by the Agency.

#### SECTION VI. INQUIRIES AND RESPONSES TO THIS NOTICE

If you have any questions regarding the requirements and procedures established by this Notice, call the contact person(s) listed in Attachment A, the <u>Data Call-In Chemical Status</u> Sheet.

All responses to this Notice (other than voluntary cancellation requests and generic data exemption claims) must include a completed <u>Data Call-In Response Form</u> and a completed <u>Requirements Status and Registrant's Response Form</u> (Attachment B for generic data and Attachment C for product specific data) and any other documents required by this Notice, and should be submitted to the contact person(s) identified in Attachment A. If the voluntary cancellation or generic data exemption option is chosen, only the <u>Data Call-In Response Form</u> need be submitted.



The Office of Compliance Monitoring (OCM) of the Office of Pesticides and Toxic Substances (OPTS), EPA, will be monitoring the data being generated in response to this Notice.

Sincerely yours,

Peter P. Caulkins Ph.D., Acting Director

Special Review and

Peter Caullans

Reregistration Division

#### Attachments

- A- Data Call-In Chemical Status Sheet
- B- Product-Specific Data Call-In Response Form
- C- Requirements Status and Registrant's Response Form for the Product Specific Data Call-In
- D- EPA Grouping of End-Use Products for Meeting Acute Toxicology Data
  Requirements for Reregistration
- E- EPA Acceptance Criteria
- F- List of Registrants Receiving This Notice
- G- Cost Share and Data Compensation Forms, and Product Specific Data Report Form

# Attachment A Chemical Status Sheet

## CEDARWOOD OIL DATA CALL[-]IN CHEMICAL STATUS SHEET

### INTRODUCTION

You have been sent this Product Specific Data Call-In Notice because you have product(s) containing cedarwood oil.

This <u>Product Specific Data Call-In Chemical Status Sheet</u>, contains an overview of data required by this notice, and point of contact for inquiries pertaining to the reregistration of cedarwood oil. This attachment is to be used in conjunction with (1) the Product Specific Data Call-In Notice, (2) the Product Specific Data Call-In Response Form (Attachment B), (3) the Requirements Status and Registrant's Form (Attachment C), (4) EPA's Grouping of End[-]Use Products for Meeting Acute Toxicology Data Requirement (Attachment D), (5) the EPA Acceptance Criteria (Attachment E), (6) a list of registrants receiving this DCI (Attachment F) and (7) the Cost Share and Data Compensation Forms in replying to this cedarwood oil Product Specific Data Call[-]In (Attachment G). Instructions and guidance accompany each form.

### DATA REQUIRED BY THIS NOTICE

The additional data requirements needed to complete the database for cedarwood oil are contained in the Requirements Status and Registrant's Response, Attachment C. The Agency has concluded that additional data on cedarwood oil are needed for specific products. These data are required to be submitted to the Agency within the time frame listed. These data are needed to fully complete the reregistration of all eligible cedarwood oil products.

### INQUIRIES AND RESPONSES TO THIS NOTICE

If you have any questions regarding the generic database of cedarwood oil, please contact Ron Kendall at (703) 308-8068.

If you have any questions regarding the product specific data requirements and procedures established by this Notice, please contact Frank Rubis (703) 308-8184. All responses to this Notice for the Product Specific data requirements should be submitted to:

Frank Rubis, Product Manager, Team 81
Product Reregistration Branch
Special Review and Reregistration Division 7508W
Office of Pesticide Programs
U.S. Environmental Protection Agency
Washington, D.C. 20460
RE: Cedarwood Oil



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# INSTRUCTIONS FOR COMPLETING THE DATA CALL-IN RESPONSE FORM FOR PRODUCT SPECIFIC DATA

- Item 1-4. Already completed by EPA.
- Item 5. If you wish to voluntarily cancel your product, answer "yes." If you choose this option, you will not have to provide the data required by the Data Call-In Notice and you will not have to complete any other forms. Further sale and distribution of your product after the effective date of cancellation must be in accordance with the Existing Stocks provision of the Data Call-In Notice (Section IV-C).
- Item 6. Not applicable since this form calls in product specific data only. However, if your product is identical to another product and you qualify for a data exemption, you must respond with "yes" to Item 7a (MUP) or 7B (EUP) on this form, provide the EPA registration numbers of your source(s); you would not complete the "Requirements Status and Registrant's Response" form. Examples of such products include repackaged products and Special Local Needs (Section 24c) products which are identical to federally registered products.
- Item 7a. For each manufacturing use product (MUP) for which you wish to maintain registration, you must agree to satisfy the data requirements by responding "yes."
- Item 7b. For each end use product (EUP) for which you wish to maintain registration, you must agree to satisfy the data requirements by responding "yes." If you are requesting a data waiver, answer "yes" here; in addition, on the "Requirements Status and Registrant's Response" form under Item 9, you must respond with Option 7 (Waiver Request) for each study for which you are requesting a waiver. See Item 6 with regard to identical products and data exemptions.
- Items 8-11. Self-explanatory.
- NOTE: You may provide additional information that does not fit on this form in a signed letter that accompanies this form. For example, you may wish to report that your product has already been transferred to another company or that you have already voluntarily canceled this product. For these cases, please supply all relevant details so that EPA can ensure that its records are correct.

# INSTRUCTIONS FOR COMPLETING THE REQUIREMENTS STATUS AND REGISTRANT'S RESPONSE FORM FOR PRODUCT SPECIFIC DATA

- Item 1-3 Completed by EPA. Note the unique identifier number assigned by EPA in Item 3. This number must be used in the transmittal document for any data submissions in response to this Data Call-In Notice.
- Item 4. The guideline reference numbers of studies required to support the product's continued registration are identified. These guidelines, in addition to the requirements specified in the Notice, govern the conduct of the required studies. Note that series 61 and 62 in product chemistry are now listed under 40 CFR 158.155 through 158.180, Subpart C.
- Item 5. The study title associated with the guideline reference number is identified.
- Item 6. The use pattern(s) of the pesticide associated with the product specific requirements is (are) identified. For most product specific data requirements, all use patterns are covered by the data requirements. In the case of efficacy data, the required studies only pertain to products which have the use sites and/or pests indicated.
- Item 7. The substance to be tested is identified by EPA. For product specific data, the product as formulated for sale and distribution is the test substance, except in rare cases.
- Item 8. The due date for submission of each study is identified. It is normally based on 8 months after issuance of the Reregistration Eligibility Document unless EPA determines that a longer time period is necessary.
- Item 9. Enter <u>only one</u> of the following response codes <u>for each data requirement</u> to show how you intend to comply with the data requirements listed in this table. Fuller descriptions of each option are contained in the Data Call-In Notice.
  - I will generate and submit data by the specified due date (Developing Data). By indicating that I have chosen this option, I certify that I will comply with all the requirements pertaining to the conditions for submittal of this study as outlined in the Data Call-In Notice. By the specified due date, I will also submit: (1) a completed "Certification With Respect To Data Compensation Requirements" form (EPA Form 8570-29) and (2) two completed and signed copies of the -Confidential Statement of Formula (EPA Form 8570-4).
  - 2. I have entered into an agreement with one or more registrants to develop data jointly (Cost Sharing). I am submitting a copy of this agreement. I understand that this option is available only for acute toxicity or certain efficacy data and only if EPA indicates in an attachment to this Notice that my product is similar



enough to another product to qualify for this option. I certify that another party in the agreement is committing to submit or provide the required data; if the required study is not submitted on time, my product may be subject to suspension. By the specified due date, I will also submit: (1) a completed "Certification With Respect To Data Compensation Requirements" form (EPA Form 8570-29) and (2) two completed and signed copies of the Confidential Statement of Formula (EPA Form 8570-4).

- 3. I have made offers to share in the cost to develop data (Offers to Cost Share). I understand that this option is available only for acute toxicity or certain efficacy data and only if EPA indicates in an attachment to this Data Call-In Notice that my product is similar enough to another product to qualify for this option. I am submitting evidence that I have made an offer to another registrant (who has an obligation to submit data) to share in the cost of that data. I am also submitting a completed "Certification of Offer to Cost Share in the Development Data" form. I am including a copy of my offer and proof of the other registrant's receipt of that offer. I am identifying the party which is committing to submit or provide the required data; if the required study is not submitted on time, my product may be subject to suspension. I understand that other terms under Option 3 in the Data Call-In Notice (Section III-C.1.) apply as well. By the specified due date, I will also submit: (1) a completed "Certification With Respect To Data Compensation Requirements" form (EPA Form 8570-29) and (2) two completed and signed copies of the Confidential Statement of Formula (EPA Form 8570-4).
- 4. By the specified due date, I will submit an existing study that has not been submitted previously to the Agency by anyone (Submitting an Existing Study). I certify that this study will meet all the requirements for submittal of existing data outlined in Option 4 in the Data Call-In Notice (Section III-C.1.) and will meet the attached acceptance criteria (for acute toxicity and product chemistry data). I will attach the needed supporting information along with this response. I also certify that I have determined that this study will fill the data requirement for which I have indicated this choice. By the specified due date, I will also submit a completed "Certification With Respect To Data Compensation Requirements" form (EPA Form 8570-29) to show what data compensation option I have chosen. By the specified due date, I will also submit: (1) a completed "Certification With Respect To Data Compensation Requirements" form (EPA Form 8570-29) and (2) two completed and signed copies of the Confidential Statement of Formula (EPA Form 8570-4).
- 5. By the specified due date, I will submit or cite data to upgrade a study classified by the Agency as partially acceptable and upgradable (Upgrading a Study). I will submit evidence of the Agency's review indicating that the study may be upgraded and what information is required to do so. I will provide the MRID or Accession number of the study at the due date. I understand that the conditions for this option outlined Option 5 in the Data Call-In Notice (Section III-C.1.) apply. By the specified due date, I will also submit: (1) a completed

"Certification With Respect To Data Compensation Requirements" form (EPA Form 8570-29) and (2) two completed and signed copies of the Confidential Statement of Formula (EPA Form 8570-4).

- 6. By the specified due date, I will cite an existing study that the Agency has classified as acceptable or an existing study that has been submitted but not reviewed by the Agency (Citing an Existing Study). If I am citing another registrant's study, I understand that this option is available only for acute toxicity or certain efficacy data and only if the cited study was conducted on my product, an identical product or a product which EPA has "grouped" with one or more other products for purposes of depending on the same data. I may also choose this option if I am citing my own data. In either case, I will provide the MRID or Accession number(s) for the cited data on a "Product Specific Data Report" form or in a similar format. By the specified due date, I will also submit: (1) a completed "Certification With Respect To Data Compensation Requirements" form (EPA Form 8570-29) and (2) two completed and signed copies of the Confidential Statement of Formula (EPA Form 8570-4).
- 7. I request a waiver for this study because it is inappropriate for my product (Waiver Request). I am attaching a complete justification for this request, including technical reasons, data and references to relevant EPA regulations, guidelines or policies. [Note: any supplemental data must be submitted in the format required by P.R. Notice 86-5]. I understand that this is my only opportunity to state the reasons or provide information in support of my request. If the Agency approves my waiver request, I will not be required to supply the data pursuant to Section 3(c)(2)(B) of FIFRA. If the Agency denies my waiver request, I must choose a method of meeting the data requirements of this Notice by the due date stated by this Notice. In this case, I must, within 30 days of my receipt of the Agency's written decision, submit a revised "Requirements Status and Registrant's Response" Form indicating the option chosen. I also understand that the deadline for submission of data as specified by the original data call-in notice will not change. By the specified due date, I will also submit: (1) a completed "Certification With Respect To Data Compensation Requirements" form (EPA Form 8570-29) and (2) two completed and signed copies of the Confidential Statement of Formula (EPA Form 8570-4).

Items 10-13. Self-explanatory.

NOTE: You may provide additional information that does not fit on this form in a signed letter that accompanies this form. For example, you may wish to report that your product has already been transferred to another company or that you have already voluntarily canceled this product. For these cases, please supply all relevant details so that EPA can ensure that its records are correct.

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		United	ed States Environmental	C	Protection Agency			Form Approved	
			DATA CALL-	LEJ .	NSB			OMB No. 2070-0107 2070-0057 Approval Expires 03-31-96	
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	1. Company name and Address SAMPLE COMPANY NO STREET ADDRI NO CITY, XX	Address IPANY ADDRESS XX 00000		2. Case # and Name 3150 WOO	Wood oils and gums	19	3. Date and Typ PRODUCT	3. Date and Type of DCI PRODUCT SPECIFIC	
1	4. EPA Product	5. I wish to	6. Generic Data			7. Product Specific Data	Date		-
	Registration	cancel this product regis- tration volun- terily.	6a. I am claiming a Generic Data Exemption because I obtain the active ingredient from the source EPA regis- tration number listed below.	6b. I Data i on the "Requ	6b. I agree to satisfy Generic Data requirements as indicated on the attached form entitled "Requirements Status and Registrant's Response."	7s. My product is a MUP and I agree to satisfy the MUP requirements on the attached form entitled "Requirements Status and Registrant's Response."	MUP and the MUP attached Irements nt's	7b. My product is an EUP and I agree to satisfy the EUP requirements on the attached form entitled "Requirements Status and Registrant's Response."	
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	8. Certification I certify that the statements made on this form and all atil acknowledge that any knowingly false or misleading states or both under applicable law.	atements made on the syknowingly false of the law.	tachments nent may	are true, accu be punishable t	i are true, accurate, and complete. be punishable by fine, imprisonment	6	9. Date		
•	10. Name of Company Contact	ontact				11.	11. Phone Number		1

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		Washington,	р. С	20460			***	OMB No. 2070-0107	0107
		REQUIREMENTS STATUS AND		REGISTRANT'S	RESPONSE			zoro- aproval Expi	Approval Expires 03-31-96
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	NO STREET ADDRESS NO CITY, XX 000	00000	EPA Reg	eg. No.	NNNNN - NNNNN	1 -	TATATA	- WWW	
	4. Guideline Requirement	5. Study Title	2≪0⊢0	Progress Reports	6. Use Pattern	7. Test Substance	€0 E	8. Time Frame	9. Registrant Response
	Number		00-	1 2 3					
<u> </u>		Prod Chem - Regular Chemical							
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		process					PER	\$650	
	61-2(b)	Discussion of formation of (1,3)		- 大 大 大 大	ABCDEFGHIJKLMNO MP/EP	and	TGAI	B mos.	
		impurities  Preliminary analysis  (1.4)			ABCDEFGHIJKLMNO	MP/EP and	TGAI	8 mos.	
-	62.2	its			ABCDEFGHIJKLANO	MP/EP		mog	
<u> </u>	62-3	Analytical method (1)			ABCDEFGHIJKIMNO ABCDEFGHIJKIMNO	MP/RP and	TCAT	8 mos.	
	, m	Physical state			ABCDEFGHIJKLMNO	MP/EP and	TGAI	8 mos.	
	63.4	1000			ABCDEFGHIJKLMNO	MP/EP and	TGAI	-11-11-	
	63 - 5 7 - 5	Melting point (6)			ABCDEFGHIJKIMNO ARCDEFGHTIKIMNO	TGAI		8 mos.	
	- 7		-		ABCDEFGHIJKLMNO MP/EP	MP/EP and	TGAI	MOB	
	10. Certification					11. Date			
	i certify that the statements i acknowledge that any knowing or both under applicable law.	made on this form and all attachments gly false or misleading statement may k	are true, a e punishabl	are true, accurate, and complete. Me punishable by fine, imprisonment	and complete. :, imprisonment	<del>- •-</del>	•		
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C. 20460  SISTRANT'S RESPONSE  Approvate Expression and supply the information requested on this form.  In the information requested on this form.  Substance  A BCDEFGHIJKLANNO TGAI/PAI  A BCDEFGHIJKLANNO TGAI/PAI  B MOS.		United States Environmental	Pr	on Agency		Form Approved	-
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63-9   Vapor pressure   ABCDEFGHLUXLAMO TGAL/PAI   8 mos     63-10   Cetanol/vater partition   (8)   ABCDEFGHLUXLAMO TGAL/PAI   8 mos     63-11   Cetanol/vater partition   (8)   ABCDEFGHLUXLAMO PAI   8 mos     63-12   Stebility   (19)   ABCDEFGHLUXLAMO MP/EP   8 mos     63-13   Stebility   (19)   ABCDEFGHLUXLAMO MP/EP   8 mos     63-14   Stepisos tebility   (12)   ABCDEFGHLUXLAMO MP/EP   8 mos     63-15   Stepisos tebility   (12)   ABCDEFGHLUXLAMO MP/EP   8 mos     63-16   Exploability   (13)   ABCDEFGHLUXLAMO MP/EP   8 mos     63-18   Viscos ty   (14)   ABCDEFGHLUXLAMO MP/EP   8 mos     63-19   Miscibility   (14)   ABCDEFGHLUXLAMO MP/EP   8 mos     63-20   Corrosion characteristics   (14)   ABCDEFGHLUXLAMO MP/EP   8 mos     63-21   Acute oral toxicity-rat   (1,3,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-2   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute infantion or this pase   (1,1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute infantion or this pase   (1,1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute infantion or this pase   (1,1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-3   Acute dermal   (1,2,37)   ABCDEFGHLUXLAMO MP/EP and TGAI   8 mos     81-4   Acute dermal   (1,2,	3-	solubility			GAI/PAI	444.5	
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KLM O EP 8	81-4 81-5 6 5	Primary eye irritation-rabbit (2) Primary dermal irritation (1,2) Dermal sensitization (4)  Efficacy - Invertebrate Control Agents Premises Treatments		a e e e e e e e e e e e e e e e e e e e			MP/BP MP/BP MP/BP		8 mos. 8 mos.	
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## Protection Agency 20460 Washington, D. C. States Environmental United

# POOTNOTES AND KEY DEFINATIONS FOR GUIDBLINE REQUIREMENTS

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that; EP = end-use product; provided formulators purchase their active ingredient(s) from a registered source, they need not submit or cite in the product that is product; provided formulators purchase their active ingrequents, from a registered source, they need not sugmit or city to the purchased and any use for the product does. not differ from those of the purchased and registered source, users are not subject to any data requirements identified in the tables.]; IEP = typical end-use product; IGAI = technical grade of the active ingredient; PAI = "pure" active ingredient; PAIRA = "pure" active ingredient, radiolabeled. 5.

Use Categories Key:

- Greenhous a nonfood crop D - Aquatic ford crop C - Terrestrial nonfood crop H - Greenhouse food crop B - Terrestrial food feed crop G - Aquatic nonfood residential - Aquatic nonfood Industrial A - Terrestrial food crop

E - Aquatic nonfood outdoor J - Forestry

0 - Indoor residential

L - Indoor food

FOOLINGES: [The following notes are referenced in column two (5. Study Title) of the REQUIREMENTS STATUS AND REGISTRANT'S RESPONSE form.] N - Indoor Medical M - Indoor nonfood K - Residential outdoor

## Prod Chem - Regular Chemical

- \*158.175 for certification of limits (62-2); and \*158.180 for enforcement Requirements pertaining to product identity, composition, analysis, and certification of ingredients are detailed further in the following sections: \*158.155 for product identity and composition (61-1); \*158.160, 158.162, and 158.165 for description of starting materials and manufacturing process (61-2); \*158.167 for discussion of formation of impurities (61-3); \*158.170 for preliminary analysis (62-1);
  - A schematic diagram and/or brief description of the production process will suffice if the pesticide is not already under full scale production and an experimental analytical methods (62-3).
- If the pesticide is not already under full scale production and an experimental use permit is sought, a discussion of unintentional ingredients shall be submitted to use permit is being sought.
  - If the technical grade of Active Ingredient cannot be isolated, a statement of composition of the practical equivalent of the technical grade of Active Ingredient must be submitted. To support registration of an MP or EP, whether produced by an integrated system or not, the technical grade of Active ingredient must be analyzed. Data on EPs or MPs will be required on a case-by-case basis. the extent this information is available.
    - Certified limits are not required for inert ingredients in products proposed for experimental use.
      - Required if technical chemical is solid at room temperature.
- technical chemical is liquid at room temperature. Required
  - technical chemical is organic and non-polar. Required if
    - test substances are dispersible with water. Required if
- product contains an oxidizing or reducing agent. Required
  - product contains combustible liquids. Required if
    - product is potentially explosive. Required if
      - product is a liquid. Required if
- product is an emulsifiable liquid and is to be diluted with petroleum solvents. Required if
  - end-use product is liquid and is to be used around electrical equipment. Required if

# Acute Toxic - Regular Chemical

- Not required if test material is corrosive to skin or has pH less than 2 or greater than 11.5; such a product will be classified as Toxicity Category 1 on the basis 1 Not required if test material is a gas or highly volatile. 2 Not required if test material is corrosive to skin or has a of potential eye and dermal irritation effects.

# United States Environmental Protection Agency Washington, D. C.

# POOTNOTES AND KEY DEFINATIONS FOR GUIDELINE REQUIREMENTS

Wood oils and gums Case # and Name: 3150

## Footnotes (cont.):

- 3 Required if the product consists of, or under conditions of use will result in, an inhalable material (e.g., gas, volatile substances, or aerosol/particulate).
  4 Applied unless repeated dermal exposure does not account and account of the control of the contro
  - anired unless repeated dermal exposure does not occur under conditions of use.

..ed a , tential to adversely affect the visual system. Registrants should consult with the agency for development of protocols and methodology · in (acute, subchronic, and/or chronic) is required for organophospates, and may be required for other cholinesterase inhibitors and other pesticides

applicators specified in 40 CFR 152.170(b) or the criteria for initiation of special review specified in 40 CFR 154.7 (a)(1).

# Efficacy - Invertebrate Control Agents

- develop and maintain the relevant data upon which the determination of efficacy is based. The Agency reserves the right to require, on a case-by-case basis (e.g., through testing that his products are efficacious when used in accordance with label directions and commenty accepted pest control practices. The registrant must significant new uses or benefits data in cases of special reviews) submission of efficacy data for any pesticide product, registered or proposed for registration The agency has waived all requirements to submit efficacy data for invertebrate control agents for nonpublic health uses. Mowever, each registrant must ensure when necessary.

  - Efficacy evaluations can be conducted under laboratory, greenhouse, or field conditions. Required to be developed and maintained in the Reggistrant's file for all pests claimed on the label when resistance to the pestcide has been demonstrated.

#### Attachment D

EPA Grouping of End-Use Products for Meeting Data Requirements for Reregistration

#### EPA'S DECISION ON BATCHING PRODUCTS CONTAINING CEDAR WOOD OIL FOR PURPOSES OF MEETING ACUTE TOXICITY DATA REQUIREMENTS FOR REREGISTRATION

In an effort to reduce the time, resources and number of animals needed to fulfill the acute toxicity data requirements for reregistration of products containing the active ingredient cedar wood oil, the Agency considered batching products. This process involves grouping similar products for purposes of acute toxicity. Factors considered in the sorting process include each product's active and inert ingredients (identity, percent composition and biological activity), type of formulation (e.g., emulsifiable concentrate, aerosol, wettable powder, granular, etc.), and labeling (e.g., signal word, use classification, precautionary labeling, etc.). Note that the Agency is not describing batched products as "substantially similar" since some products within a batch may not be considered chemically similar or have identical use patterns.

Batching has been accomplished using the information described above as available. Acute toxicity data on individual products has frequently been found to be incomplete. Notwithstanding the batching process, the Agency reserves the right to require, at any time, acute toxicity data for an individual product should the need arise.

Registrants of products within a batch may choose to cooperatively generate, submit or cite a single battery of six acute toxicological studies to represent all the products within that batch. It is the registrants' option to participate in the process with all other registrants, only some of the other registrants, or only their own products within a batch, or to generate all the required acute toxicological studies for each of their own products. If a registrant chooses to rely upon previously submitted acute toxicity data, he/she may do so provided that the data base is complete and valid by today's standards (see acceptance criteria attached), the formulation tested is considered by EPA to be similar for acute toxicity, and the formulation has not been significantly altered since submission and acceptance of the acute toxicity data. Regardless of whether new data is generated or existing data is cited, the registrant must clearly identify the material tested by its EPA registration number.

In deciding how to meet the product specific data requirements, registrants must follow the directions given in the Data Call-In Notice and its attachments appended to the RED. The DCI Notice contains two response forms which are to be completed and submitted to the Agency within 90 days of receipt. The first form, "Data Call-In Response", asks whether the registrant will meet the data requirements for each product. The second form, "Requirements Status and Registrant's Response", lists the product specific data required for each product, including the standard six acute toxicity tests. A registrant who wishes to participate in a batch must decide whether he/she will provide the data or depend on someone else to do so. If a registrant supplies the data to support a batch of products, he/she must select one of the following options: Developing Data (Option 1), Submitting an Existing Study (Option 4), Upgrading an Existing Study (Option 5), or Citing an Existing Study (Option 6). If a registrant depends on another's data, he/she-must-choose among: Cost Sharing (Option 2), Offers to Cost Share (Option 3) or Citing an Existing Study (Option 6). If a registrant does not want to participate in a batch, the choices are Options 1, 4, 5 or 6. However, a registrant should know that choosing not to participate in a batch does not preclude other registrants in the batch form citing his/her studies and offering to cost share (Option 3) those studies.



Table I lists the products of Batch 1.

Table I.

Batch No.	EPA Reg. No.	% of Cedar Wood Oil	Formulation Type
1	65555-1	5.17	block
	. 65813-1	4.40	block
	66211-1	4.40	block

Table II lists the products which could not be batched. These products were not considered similar for purposes of acute toxicity.

The registrants of these products are responsible for meeting the acute toxicity data requirements specified in the data matrix for end-use products.

Table II.

EPA Reg. No.	% of Cedar Wood Oil & other Active Ingredients	Formulation Type
63380-1	0.48	aerosol
42443-1	0.50 Oil of Pennyroyal 2.00 Oil of Eucalyptus 1.00 Oil of Citronella 0.50 Oil of Rue 0.12	collar

### ATTACHMENT E EPA ACCEPTANCE CRITERIA

#### SUBDIVISION D

Guideline	Study Title
Series 61	Product Identity and Composition
Series 62	Analysis and Certification of Product Ingredients
Series 63	Physical and Chemical Characteristics

#### 61 Product Identity and Composition

#### ACCEPTANCE CRITERIA

Does yo	our study meet the following acceptance criteria?
1	Name of technical material tested (include product name and trade name, if appropriate).
2	Name, nominal concentration, and certified limits (upper and lower) for each active ingredient and each intentionally-added inert ingredient.
3	Name and upper certified limit for each impurity or each group of impurities present at $\geq 0.1\%$ by weigh and for certain toxicologically significant impurities (e.g., dioxins, nitrosamines) present at $< 0.1\%$ .
4	Purpose of each active ingredient and each intentionally-added inert.
5	Chemical name from Chemical Abstracts index of Nomenclature and Chemical Abstracts Service (CAS Registry Number for each active ingredient and, if available, for each intentionally-added inert.
6	Molecular, structural, and empirical formulas, molecular weight or weight range, and any company assigned experimental or internal code numbers for each active ingredient.
7	Description of each beginning material in the manufacturing process.  EPA Registration Number if registered; for other beginning materials, the following:  Name and address of manufacturer or supplier.  Brand name, trade name or commercial designation.  Technical specifications or data sheets by which manufacturer or supplier describes composition properties or toxicity.
8I	Description of manufacturing process.  Statement of whether batch or continuous process.

	Relative amounts of beginning materials and order in which they are added.
	Description of equipment.
	Description of physical conditions (temperature, pressure, humidity) controlled in each step and the parameters that are maintained.
	Statement of whether process involves intended chemical reactions.
	Flow chart with chemical equations for each intended chemical reaction.
	Duration of each step of process.
	Description of purification procedures.
	Description of measures taken to assure quality of final product.
)	Discussion of formation of impurities based on established chemical theory addressing (1) each impurity which may be present at $\geq 0.1\%$ or was found at $\geq 0.1\%$ by product analyses and (2) certain toxicologically significant impurities (see #3).

#### 62 Analysis and Certification of Product Ingredients

#### ACCEPTANCE CRITERIA

The following criteria apply to the technical grade of the active ingredient being reregistered. Use a table to present the information in items 6, 7, and 8.

Does your study meet the following acceptance criteria? Five or more representative samples (batches in case of batch process) analyzed for each active ingredient and all impurities present at  $\geq 0.1\%$ . Degree of accountability or closure  $\geq ca$  98%. 2.\_\_\_\_ Analyses conducted for certain trace toxic impurities at lower than 0.1% (examples, nitrosamines in the case of products containing dinitroanilines or containing secondary or tertiary amines/alkanolamines plus nitrites; polyhalogenated dibenzodioxins and dibenzofurans). [Note that in the case of nitrosamines both fresh and stored samples must be analyzed.1. Complete and detailed description of each step in analytical method used to analyze above samples. Statement of precision and accuracy of analytical method used to analyze above Identities and quantities (including mean and standard deviation) provided for each 6.\_\_\_\_ analyzed ingredient. Upper and lower certified limits proposed for each active ingredient and intentionally 7. added inert along with explanation of how the limits were determined. Upper certified limit proposed for each impurity present at  $\geq 0.1\%$  and for certain 8. toxicologically significant impurities at <0.1% along with explanation of how limit determined. 9. \_\_\_ Analytical methods to verify certified limits of each active ingredient and impurities (latter not required if exempt from requirement of tolerance or if generally recognized as safe by FDA) are fully described. Analytical methods (as discussed in #9) to verify certified limits validated as to their 10.



precision and accuracy.

#### 63 Physical and Chemical Characteristics

#### ACCEPTANCE CRITERIA

The following criteria apply to the technical grade of the active ingredient being reregistered.  Does your study meet the following acceptance criteria?		
Verbal description of coloration (or lack of it)  Any intentional coloration also reported in terms of Munsell color system		
63-3 Physical State		
Verbal description of physical state provided using terms such as "solid, granular, volatile liquid"  Based on visual inspection at about 20-25° C		
63-4 Odor		
Verbal description of odor (or lack of it) using terms such as "garlic-like, characteristic of aromatic compounds"		
Observed at room temperature		
63-5 Melting Point		
Reported in °C		
Any observed decomposition reported		
63-6 Boiling Point		
Reported in °C		
Pressure under which B.P. measured reported		
Any observed decomposition reported		
63-7 Density, Bulk Density, Specific Gravity Measured at about 20-25° C		
Density of technical grade active ingredient reported in g/ml or the specific gravity of liquids reported with reference to water at 20° C. [Note: <u>Bulk</u> density of registered products may be reported in lbs/ft <sup>3</sup> of lbs/gallon.]		
63-8 Solubility		
Determined in distilled water and representative polar and non-polar solvents, including those used formulations and analytical methods for the pesticide  Measured at about 20-25° C		
Reported in g/100 ml (other units like ppm acceptable if sparingly soluble)		

53-9 Vapo	or Pressure	
	Measured at 25° C (or calculated by extrapolation from measurements made at higher temperature pressure too low to measure at 25° C)	j
	Experimental procedure described	
	Reported in mm Hg (torr) or other conventional units	
63-10 Dis	sociation Constant	
	Experimental method described	
	Temperature of measurement specified (preferably about	
	20-25°C)	
63-11 Oct	tanol/water Partition Coefficient	
	Measured at about 20-25° C	
	Experimentally determined and description of procedure provided (preferred method-45 Fed. Regis 77350)	te
	Data supporting reported value provided	
63-12 pH		
	Measured at about 20-25° C	
	Measured following dilution or dispersion in distilled water	
63-13 Sta	bility	
	Sensitivity to metal ions and metal determined	
	Stability at normal and elevated temperatures	
	Sensitivity to sunlight determined	

#### SUBDIVISION F

<u>Guideline</u>	Study Title
81-1	Acute Oral Toxicity in the Rat
81-2	Acute Dermal Toxicity in the Rat, Rabbit or Guinea Pig
81-3	Acute Inhalation Toxicity in the Rat
81-4	Primary Eye Irritation in the Rabbit
81-5	Primary Dermal Irritation Study
81-6	Dermal Sensitization in the Guinea Pig

#### 81-1 Acute Oral Toxicity in the Rat

#### ACCEPTANCE CRITERIA

1 Identify material tested (technical, end-use product, etc).	•
2. At least 5 young adult rats/sex/group.	•
3. Dosing, single oral may be administered over 24 hrs.	
4. Vehicle control if other than water.	
5. Doses tested, sufficient to determine a toxicity category or a limit dose (5000	mg/kg).
6. Individual observations at least once a day.	
7. Observation period to last at least 14 days, or until all test animals appear normal	whichever
is longer.	
8 Individual daily observations.	
9. Individual body weights.	
10 Gross necropsy on all animals.	

#### 81-2 Acute Dermal toxicity in the Rat, Rabbit or Guinea Pig

#### ACCEPTANCE CRITERIA

1.	Identify material tested (technical, end-use product, etc).
2.	At least 5 animals/sex/group.
3. <del>*</del>	Rats 200-300 gm, rabbits 2.0-3.0 kg or guinea pigs 350-450 gm.
4.	Dosing, single dermal.
5.	Dosing duration at least 24 hours.
6. <u>*</u>	Vehicle control, only if toxicity of vehicle is unknown.
7.	Doses tested, sufficient to determine a toxicity category or a limit dose (2000 mg/kg).
8.	Application site clipped or shaved at least 24 hours before dosing.
9.	Application site at least 10% of body surface area.
0.	Application site covered with a porous nonirritating cover to retain test material and
re	vent ingestion.
1.	Individual observations at least once a day.
2.	Observation period to last at least 14 days.
3.	<del></del>
4.	Gross necropsy on all animals.

#### 81-3 Acute Inhalation Toxicity in the Rat

#### ACCEPTANCE CRITERIA

<ul> <li>Product is a gas, a solid which may produce a significant vapor hazard based on toxicity and expect use or contains particles of inhalable size for man (aerodynamic diameter 15 μm or less).</li> <li>At least 5 young adult rats/sex/group.</li> <li>Dosing, at least 4 hours by inhalation.</li> <li>Chamber air flow dynamic, at least 10 air changes/hour, at least 19% oxygen content.</li> <li>Chamber temperature, 22° C (±2°), relative humidity 40-60%.</li> <li>Monitor rate of air flow.</li> <li>Monitor actual concentrations of test material in breathing zone.</li> <li>Monitor aerodynamic particle size for aerosols.</li> <li>Doses tested, sufficient to determine a toxicity category or a limit dose (5 mg/L actual concentrations)</li> </ul>	rted
At least 5 young adult rats/sex/group.  Dosing, at least 4 hours by inhalation.  Chamber air flow dynamic, at least 10 air changes/hour, at least 19% oxygen content.  Chamber temperature, 22° C (±2°), relative humidity 40-60%.  Monitor rate of air flow.  Monitor actual concentrations of test material in breathing zone.  Monitor aerodynamic particle size for aerosols.	
Dosing, at least 4 hours by inhalation.  Chamber air flow dynamic, at least 10 air changes/hour, at least 19% oxygen content.  Chamber temperature, 22° C (±2°), relative humidity 40-60%.  Monitor rate of air flow.  Monitor actual concentrations of test material in breathing zone.  Monitor aerodynamic particle size for aerosols.	
6. Chamber temperature, 22° C (+2°), relative humidity 40-60%.  7. Monitor rate of air flow.  8. Monitor actual concentrations of test material in breathing zone.  9. Monitor aerodynamic particle size for aerosols.	
6. Chamber temperature, 22° C (+2°), relative humidity 40-60%.  7. Monitor rate of air flow.  8. Monitor actual concentrations of test material in breathing zone.  9. Monitor aerodynamic particle size for aerosols.	
6. Chamber temperature, 22° C (+2°), relative humidity 40-60%.  7. Monitor rate of air flow.  8. Monitor actual concentrations of test material in breathing zone.  9. Monitor aerodynamic particle size for aerosols.	
7. Monitor rate of air flow. 8. Monitor actual concentrations of test material in breathing zone. 9. Monitor aerodynamic particle size for aerosols. 10. Doses tested, sufficient to determine a toxicity category or a limit dose (5 mg/L actual concentral).	
8. Monitor actual concentrations of test material in breathing zone.  9. Monitor aerodynamic particle size for aerosols.  10. Doses tested sufficient to determine a toxicity category or a limit dose (5 mg/L actual concentration).	
9. Monitor aerodynamic particle size for aerosols.  10. Doses tested, sufficient to determine a toxicity category or a limit dose (5 mg/L actual concentral).	
10 Doses tested sufficient to determine a toxicity category or a limit dose (5 mg/L actual concentra	
10 Dobob tostod, carriedont to determine a terment, the got, of a minute of the control of	ion
of respirable substance).	
11 Individual observations at least once a day.	•
12. Observation period to last at least 14 days.	
13 Individual body weights.	
14. Gross necropsy on all animals.	

#### 81-4 Primary Eye Irritation in the Rabbit

#### ACCEPTANCE CRITERIA

1.	Identify material tested (technical, end-use product, etc).
2.	Study not required if material is corrosive, causes severe dermal irritation or has a pH of $\leq 2$ or
	≥11.5.
3.	6 adult rabbits.
4.	Dosing, instillation into the conjunctival sac of one eye per animal.
5.	Dose, 0.1 ml if a liquid; 0.1 ml or not more than 100 mg if a solid, paste or particulate substance.
6	Solid or granular test material ground to a fine dust.
7.	Eyes not washed for at least 24 hours.
8	Eyes examined and graded for irritation before dosing and at 1, 24, 48 and 72 hr, then daily until eyes
	are normal or 21 days (whichever is shorter).
9. <u>*</u>	Individual daily observations.

#### 81-5 Primary Dermal Irritation Study

#### ACCEPTANCE CRITERIA

1.	Identify material tested (technical, end-use product, etc).
2	Study not required if material is corrosive or has a pH of $\leq 2$ or $\geq 11.5$ .
3	6 adult animals.
4.	Dosing, single dermal.
5.	Dosing duration 4 hours.
6	Application site shaved or clipped at least 24 hours prior to dosing.
7.	Application site approximately 6 cm <sup>2</sup> .
8.	Application site covered with a gauze patch held in place with nonirritating tape.
9.	Material removed, washed with water, without trauma to application site.
10.	Application site examined and graded for irritation at 1, 24, 48 and 72 hr, then daily until normal o
	14 days (whichever is shorter).
11. <u>*</u>	Individual daily observations.

#### 81-6 Dermal Sensitization in the Guinea Pig

#### ACCEPTANCE CRITERIA

## ATTACHMENT F LIST OF ALL REGISTRANTS SENT THIS DATA CALL-IN NOTICE

### ATTACHMENT G COST SHARE AND DATA COMPENSATION FORMS



#### United States Environmental Protection Agency Washington, DC 20460

#### CERTIFICATION OF OFFER TO COST SHARE IN THE DEVELOPMENT OF DATA Approval Expires 12-31-92

Form Approved OMB No. 2070-0108

Public reporting burden for this collection of information is estimated to average 15 minutes per response, including time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding the burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to Chief, Information Policy Branch, PM-223, U.S. Environmental Protection Agency, 401 M St., S.W., Washington, DC 20460; and to the Office of Management and Budget, Paperwork Reduction Project (2070-0106), Washington, DC 20503.

Piease fill in blanks below. Company Number Company Name **EPA Chemical Number** Chemical Name I Certify that: My company is willing to develop and submit the data required by EPA under the authority of the Federal Insecticide, Fungicide and Rodenticide Act (FIFRA), if necessary. However, my company would prefer to enter into an agreement with one or more registrants to develop jointly or share in the cost of developing data. My firm has offered in writing to enter into such an agreement. That offer was irrevocable and included an offer to be bound by arbitration decision under section 3(c)(2)(B)(iii) of FIFRA if final agreement on all terms could not be reached otherwise. This offer was made to the following firm(s) on the following date(s): Date of Offer Name of Firm(s) Certification: I certify that I am duly authorized to represent the company named above, and that the statements that I have made on this form and all attachments therein are true, accurate, and complete. I acknowledge that any knowingly false or misleading statement may be punishable by fine or imprisonment or both under applicable law. Signature of Company's Authorized Representative Date Name and Title (Please Type or Print)



#### United States Environmental Protection Agency Washington, D.C 20460

#### CERTIFICATION WITH RESPECT TO DATA COMPENSATION REQUIREMENTS

Form Approved
OMB No. 2070-0106

Approval Expires 12-31-92

Public reporting burden for this collection of information is estimated to average 15 minutes per response, including time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding the burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to Chief, Information Policy Branch, PM-223, U.S. Environmental Protection Agency, 401 M. St., S.W., Washington, D.C. 20460; and to the Office of Management and Budget, Paperwork Reduction Project (2070-0106), Washington, D.C. 20503.

Paperwork Reduction Project (2070-0106), Washington, D.C. 20503.	
Instructions	
Please fill in Manks below.	•
Company Name	Company Number
Chemical Name	EPA Chemical Number
I Certify that:	•
1. For each study cited in support of reregistration under the Federal Is an exclusive use study, I am the original data submitter, or I have obtained that study.	assecticide, Fungicide and Rodenticide Act (FIFRA) that is alined the written permission of the original data submitter to
2. That for each study cited in support of reregistration under FIFRA submitter, or I have obtained the written permission of the original data that submitted data I have cited and have offered to: (a) Pay compensation(2)(D) of FIFRA; and (b) Commence negotiation to determine where FIFRA and the amount of compensation due, if any. The companies I	ta subminer, or I have notified in writing the company(ics) tion for those data in accordance with section 3(c)(1)(D) and ich data are subject to the compensation requirement of
[ ] All companies on the data submitters' list for he active ing Method or Cite-All option under the Selective Method). (Also	redient listed on this form (Cite-All sign the General Offer to Pay below.)
[ ] The companies who have submitted the studies listed on the indicated on the attached "Requirements Status and Registrants relying in support of reregistration. (Selective Method)	
<ol> <li>That I have previously complied with section 3(c)(1)(D) of FIFRA under FIFRA.</li> </ol>	for the studies I have cited in support of reregistration
Signature	Date
Name and Title (Please Print)	
GENERAL OFFER TO PAY: I hereby offer and agree to pay comper my products, to the extent required by FIFRA section 3(c)(1)(D) and 3(	
Signature	Date
Name and Title (Please Print)	



